

### Training on Galaxy: Metagenomics February 2017

## Find Rapidly OTU with Galaxy Solution

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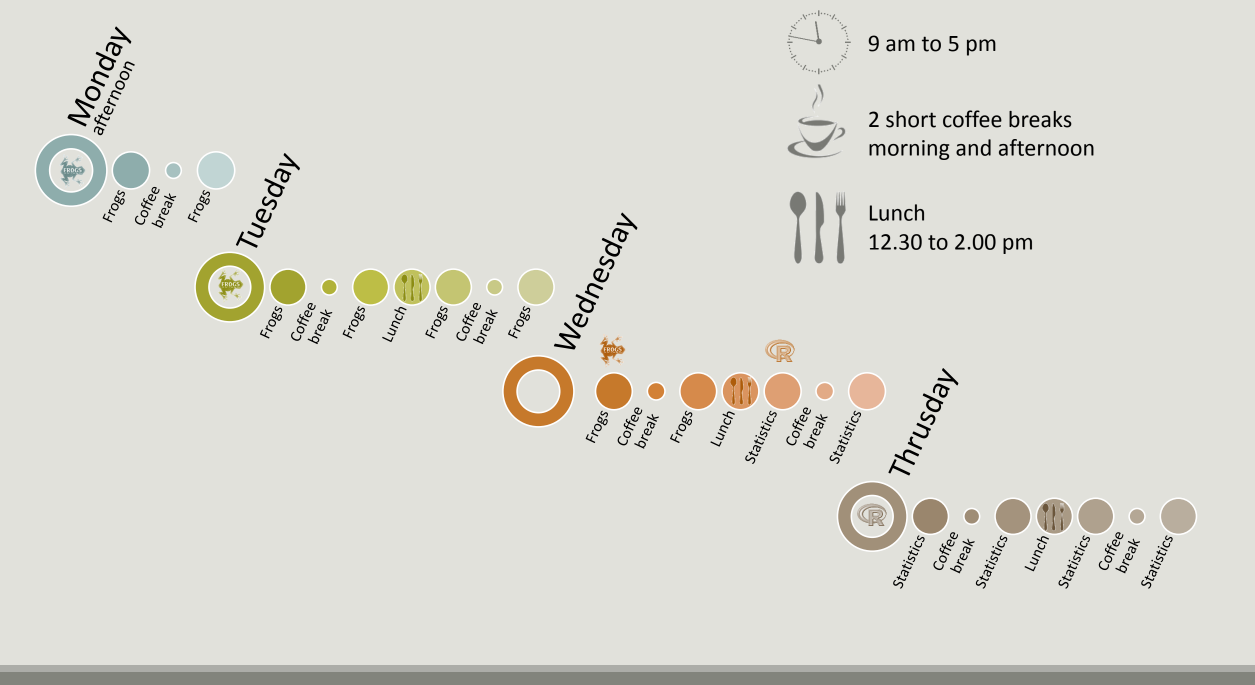
\*THESE AUTHORS HAVE CONTRIBUTED EQUALLY TO THE PRESENT WORK.



### Feedback:

### What are your needs in "metagenomics"?

Your background ?

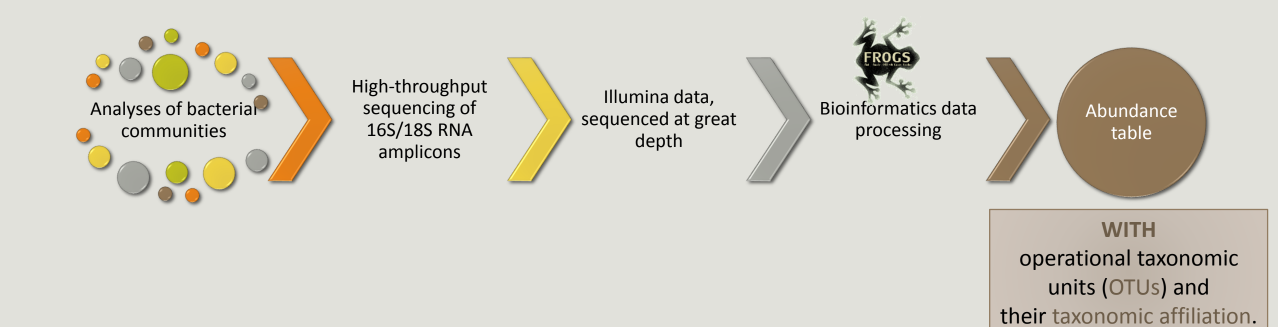




- Objectives
- Material: data + FROGS
- Data upload into galaxy environment
- Demultiplex tool
- Preprocessing
- Clustering + Cluster Statistics
- Chimera removal

- Filtering
- Affiliation + Affiliation Statistics
- Normalization
- Tool descriptions
- Format transformation
- Workflow creation
- Download data
- Some figures

### Objectives



### OTUs for ecology

#### **Operational Taxonomy Unit:**

a grouping of similar sequences that can be treated as a single « species »

#### Strengths:

- Conceptually simple
- Mask effect of poor quality data
  - Sequencing error
  - In vitro recombination (chimera)

#### Weaknesses:

- Limited resolution
- Logically inconsistent definition

### Objectives

	Affiliation	Sample 1	Sample 2	Sample 3	Sample 4	Sample 5	Sample 6
OTU1	Species A	0	100	0	45	75	18645
OTU2	Species B	741	0	456	4421	1255	23
OTU3	Species C	12786	45	3	0	0	0
OTU4	Species D	127	4534	80	456	756	108
OTU5	Species E	8766	7578	56	0	0	200

### Why we have developed FROGS

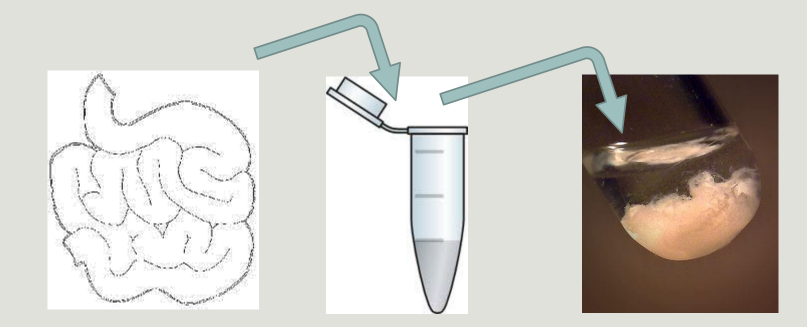
The current processing pipelines struggle to run in a reasonable time.

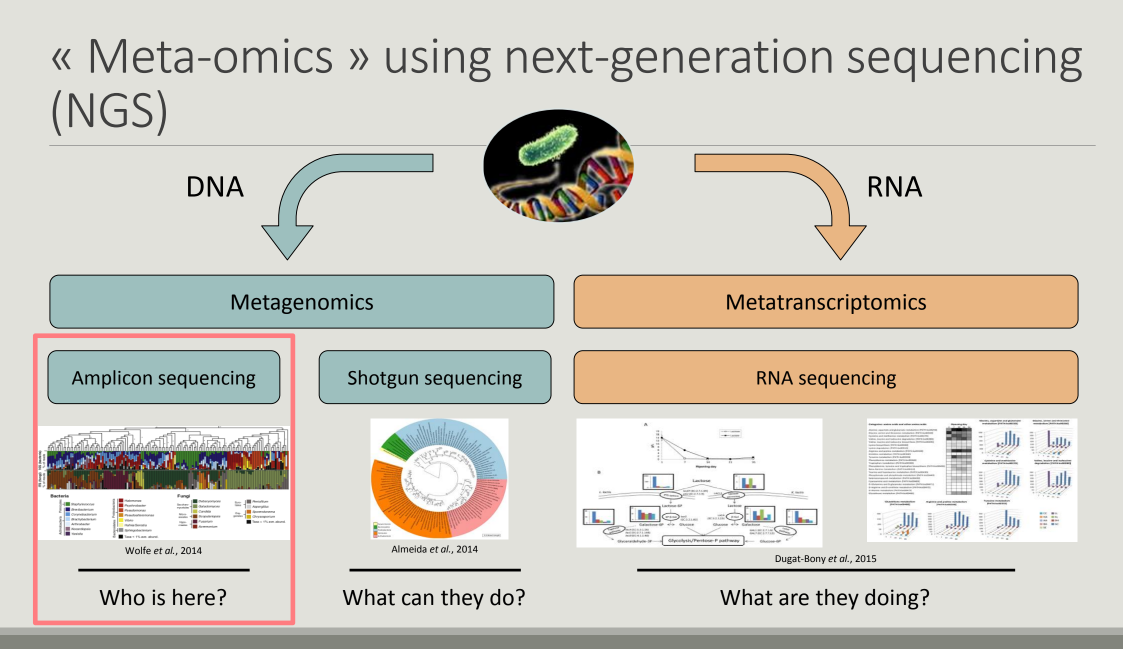
The most effective solutions are often designed for specialists making access difficult for the whole community.

In this context we developed the pipeline FROGS: « Find Rapidly OTU with Galaxy Solution ».

## Material

### Sample collection and DNA extraction





## The gene encoding the small subunit of the ribosomal RNA

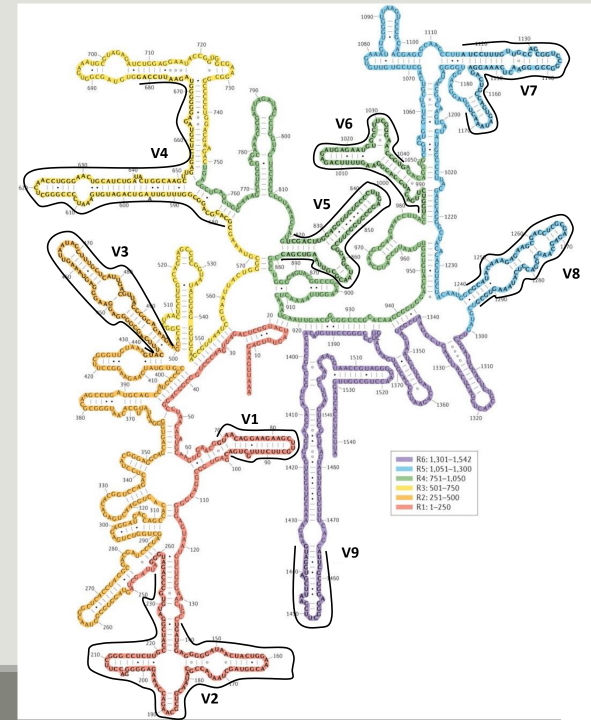
The most widely used gene in **molecular phylogenetic** studies

Ubiquist gene : 16S rDNA in prokayotes ; 18S rDNA in eukaryotes

Gene encoding a ribosomal RNA : non-coding RNA (not translated), part of the small subunit of the ribosome which is responsible for the translation of mRNA in proteins

Not submitted to lateral gene transfer

Availability of databases facilitating comparison (Silva 2015: >22000 type strains)



### Secondary structure of the 16S rRNA of

#### Escherichia coli

In red, fragment R1 including regions V1 and V2; in orange, fragment R2 including region V3; in yellow, fragment R3 including region V4; in green, fragment R4 including regions V5 and V6; in blue, fragment R5 including regions V7 and

V8;

and in purple, fragment R6 including region V9.

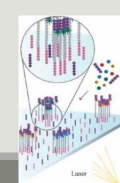
Uniting the classification of cultured and uncultured bacteria and archaea using 16S rRNA gene sequences Pablo Yarza, et al. Nature Reviews Microbiology 12, 635–645 (2014) doi:10.1038/nrmicro3330

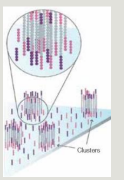
## The gene encoding the small subunit of the ribosomal RNA



### Steps for Illumina sequencing

- 1st step : one PCR
   2nd step: one PCR
   2nd step: one PCR
- 3<sup>rd</sup> step: on flow cell, the cluster generations
- 4<sup>th</sup> step: sequencing





#### Amplification and sequencing

« Universal » primer sets are used for PCR amplification of the phylogenetic biomarker

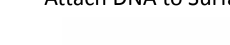
The primers contain adapters used for the sequencing step and barcodes (= tags = MIDs) to distinguish the samples (multiplexing = sequencing several samples on the same run)

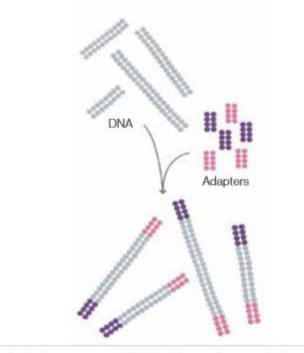


### Cluster generation

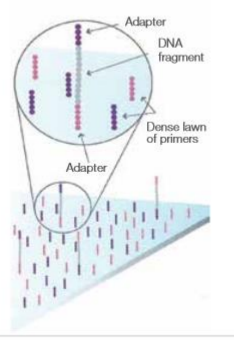
Prepare Genomic DNA Sample

Attach DNA to Surface



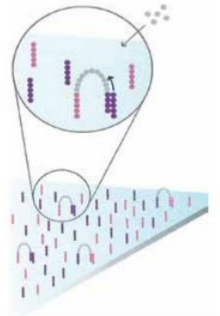


Randomly fragment genomic DNA and ligate adapters to both ends of the fragments.



Bind single-stranded fragments randomly to the inside surface of the flow cell channels.

Attach DNA to surface



Bridge Amplification

Add unlabeled nucleotides and enzyme to initiate solid-phase bridge amplification.

Bridge amplification

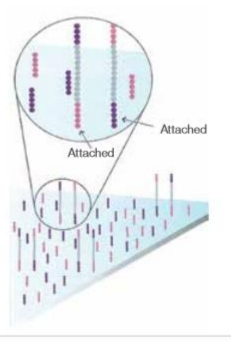
### Cluster generation

#### Fragments Become Double Stranded Denature the Double-Stranded Molecules

## Attached Attached Free terminus

The enzyme incorporates nucleotides to build double-stranded bridges on the solid-phase substrate.

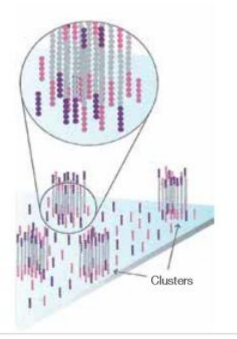
Fragments become double stranded



Denaturation leaves single-stranded templates anchored to the substrate.

Denature the double-stranded molecule

#### Complete Amplification



Several million dense clusters of double-stranded DNA are generated in each channel of the flow cell.

Cycle of new strand synthesis and denaturation to make multiple copies of the same sequence (amplification) Reverse strands are washed

### Sequencing by synthesis

**Determine First Base** 

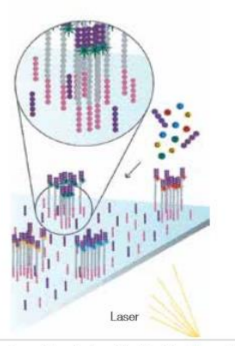
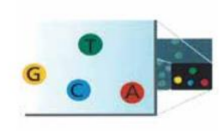
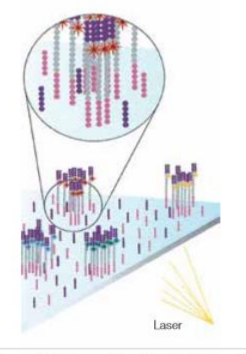


Image First Base



Determine Second Base



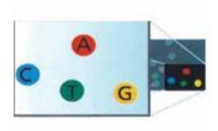
The first sequencing cycle begins by adding four labeled reversible terminators, primers, and DNA polymerase.

Light signal is more strong in cluster

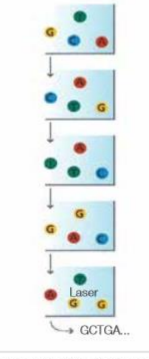
After laser excitation, the emitted fluorescence from each cluster is captured and the first base is identified. The next cycle repeats the incorporation of four labeled reversible terminators, primers, and DNA polymerase.

### Sequencing by synthesis

#### Image Second Chemistry Cycle



Sequencing Over Multiple Chemistry Cycles

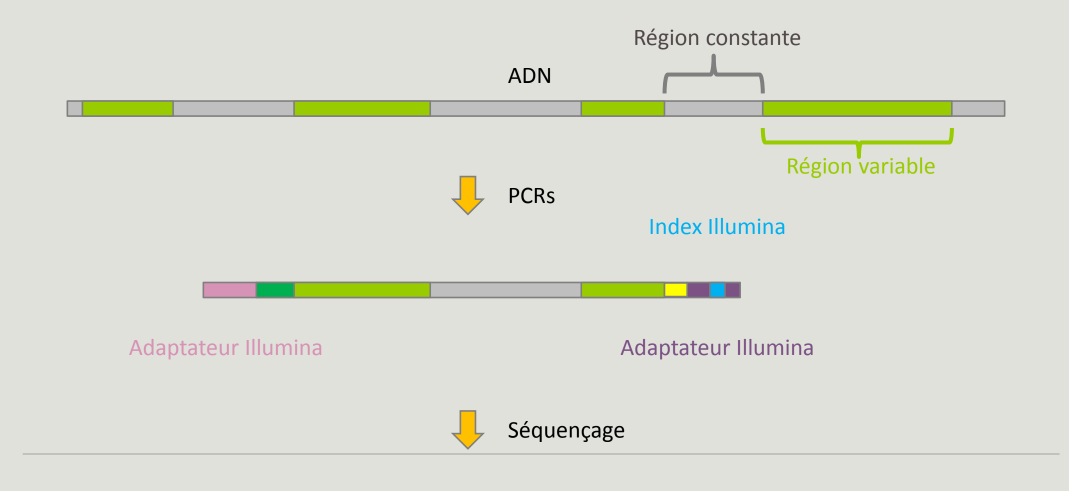


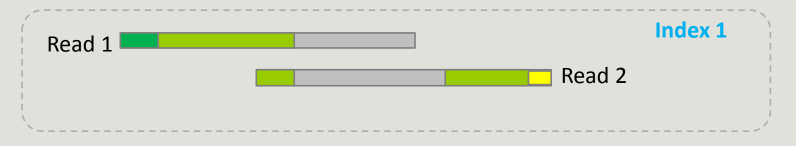
After laser excitation, the image is captured as before, and the identity of the second base is recorded.

The sequencing cycles are repeated to determine the sequence of bases in a fragment, one base at a time.

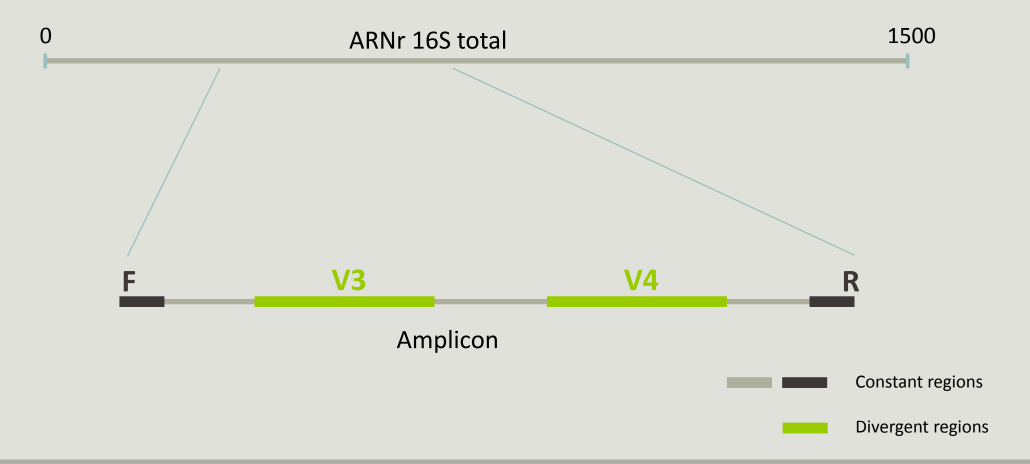
Barcode is read, so cluster is identified.

After first sequencing (250 or 300 nt of Reverse strand), fragment form bridges again and Forward strand can be sequenced also.





# Identification of bacterial populations may be not discriminating



#### Amplification and sequencing

Sequencing is generally perform on Roche-454 or Illumina MiSeq platforms.

Roche-454 generally produce ~ 10 000 reads per sample

MiSeq ~ 30 000 reads per sample

Sequence length is **>650 bp** for pyrosequencing technology (Roche-454) and **2 x 300 bp** for the MiSeq technology in paired-end mode.

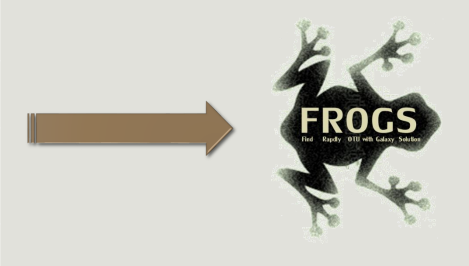


## Methods



### Which bioinformatics solutions ?

	Disadvantages
QIIME	Installation problem Command lines
UPARSE	Global clustering command lines
MOTHUR	Not MiSeq data without normalization Global hierarchical clustering Command lines
MG-RAST	No modularity No transparence



QIIME allows analysis of high-throughput community sequencing data J Gregory Caporaso et al, Nature Methods, 2010; doi:10.1038/nmeth.f.303 Introducing mothur: Open-source, platform-independent, community-supported software for describing and comparing microbial communities. Schloss, P.D., et al., Appl Environ Microbiol, 2009, doi: 10.1128/AEM.01541-09 UPARSE: Highly accurate OTU sequences from microbial amplicon reads Edgar, R.C. et al, *Nature Methods*, 2013, dx.doi.org/10.1038/nmeth.2604 The metagenomics RAST server – a public resource for the automatic phylogenetic and functional analysis of metagenomes F Meyer et al, BMC Bioinformatics, 2008, doi:10.1186/1471-2105-9-386

### FROGS ?

Use platform Galaxy

Set of modules = Tools to analyze your "big" data

Independent modules

Run on Illumina/454 data 16S, 18S, and 23S

New clustering method

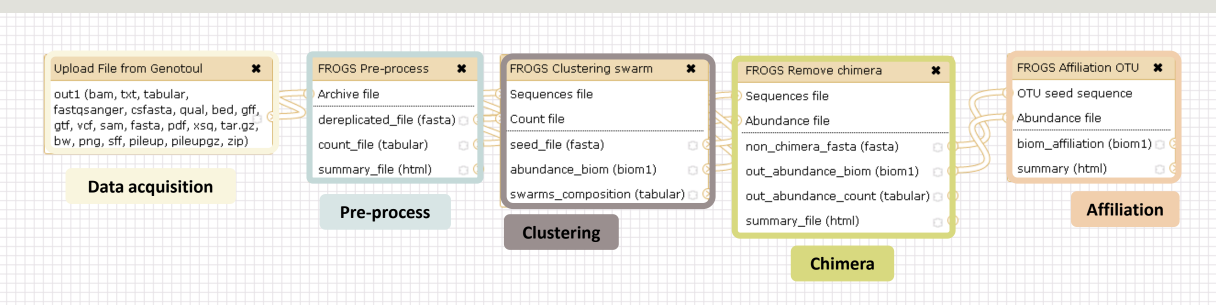
Many graphics for interpretation

User friendly, hiding bioinformatics infrastructure/complexity

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Tools	FROGS Pre-process Illumina (version 1.0.0)	🔒 History 🛛 🕹 🗘
FROGS - FIND RAPIDLY OTU WITH GALAXY SOLUTION FROGS pipeline	Input type:     Files by samples •	Unnamed history 5.0 GB
Upload archive from your computer	Samples files can be provided in single archive or with two files (R1 and R2) by sample. Reads already contiged ?: No	③19: FROGS Filters: ● ℓ X abundance table.biom
<u>Demultiplex reads</u> Split by samples the reads in function of inner barcode.	The inputs contains 1 file by sample : Reads 1 and Reads 2 are already contiged by pair. Samples	© <u>18: FROGS Filters:</u> ● ℓ ¤ summary.html
FROGS Pre-process Illumina Step 1 in metagenomics analysis from Illumina	Samples 1 Name:	Seed.fasta IT: FROGS Filters: ● Ø X Seed.fasta
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FROGS Clustering swarm Step 2 in metagenomics analysis : clustering.	Reads 1:	③15: FROGS Filters: ● Ø ⋈ abundance_table.tsv
FROGS Remove chimera Remove PCR chimera in each sample.	reads 2:	14: FROGS Clusters ● ℓ × stat: summary.html
FROGS Affiliation otu 165 Step 3 in metagenomics	R2 FASTQ file of paired-end reads.	13: FROGS Clusters ● ℓ ☆ stat: summary.html
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FROGS abundance normalisation Step 4 in metagenomics analysis	The read1 size.	★ 11: FROGS Affiliation ● Ø ※ otu 16S: tax_affiliation.biom
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FROGS Filters Step in metagenomics analysis from Illumina (165/185) : Filters on Clusters/OTUs.	Expected amplicon size:	excluded data report.html 9: FROGS Remove chimera:
FROGS Clusters stat Process some metrics on clusters.	The expected size for the majority of the amplicons (with primers).  Be minimum amplicon size:	non chimera abundance.biom
FROGS BIOM to TSV Converts a BIOM file in TSV file.	The minimum size for the amplicons (with primers).	chimera: non_chimera.fasta 7: FROGS Clustering
	Maximum amplicon size:	rente de clastering o V &



### **FROGS** Pipeline

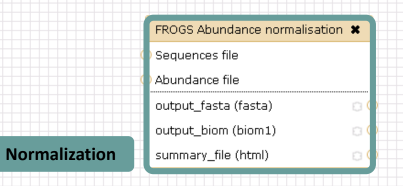




#### Upload File from Genotoul FROGS Clustering swarm FROGS Affiliation OTU × FROGS Pre-process × FROGS Remove chimera × × OTU seed sequence out1 (bam, txt, tabular, Archive file Sequences file Sequences file fastqsanger, csfasta, qual, bed, gff, Abundance file Count file dereplicated\_file (fasta) 🖂 🤇 Abundance file gtf, vcf, sam, fasta, pdf, xsq, tar.gz, bw, png, sff, pileup, pileupgz, zip) biom\_affiliation (biom1) 🖂 🤇 count\_file (tabular) seed\_file (fasta) non\_chimera\_fasta (fasta) summary (html) summary\_file (html) abundance\_biom (biom1) 00 out\_abundance\_biom (biom1) 🛛 🔅 **Data acquisition** swarms\_composition (tabular) 🗅 🤇 out\_abundance\_count (tabular) 💿 **Pre-process** summary\_file (html) Clustering Chimera

Affiliation





#### Upload File from Genotoul

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FROGS Remove chimera	×	
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out_abundance_count (tabular)	00	
summary_file (html)	00	

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OTU seed seauence	

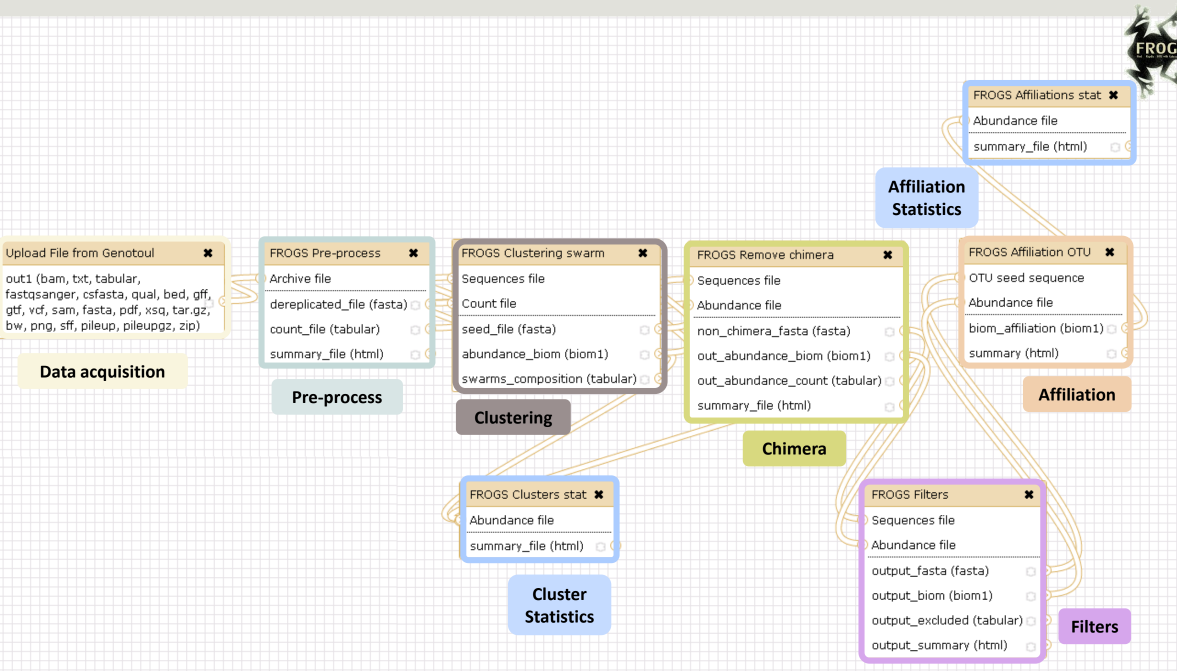
Abundance file

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summary (html)

Affiliation

#### Chimera





#### Affiliation **Statistics**

×

#### FROGS Affiliation OTU

OTU seed sequence

Abundance file

biom\_affiliation (biom1) 🖸

summary (html)

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Affiliation

**Filters** 

FROGS Filters Sequences file Abundance file output\_fasta (fasta) output\_biom (biom1) output\_excluded (tabular) 🖸 output\_summary (html)

#### Upload File from Genotoul

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#### **Data acquisition**

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**Convert to TSV** 

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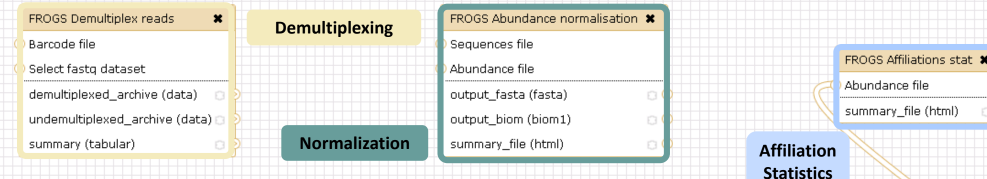
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FROGS Remove chimera

#### Chimera

FROGS TSV to BIOM X Abundance TSV File Multi\_hits TSV File biom\_file (biom1) sequence\_file (fasta) **Convert TSV to** Biom



Upload File from Genotoul	
out1 (bam, txt, tabular, fastqsanger, csfasta, qual, bed, gff, gtf, vcf, sam, fasta, pdf, xsq, tar.gz, bw, png, sff, pileup, pileupgz, zip)	

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FROGS Clusters stat 🗶 Abundance file summary\_file (html) 🛛 🔿

> Cluster **Statistics**

FROGS Remove chimera × Sequences file Abundance file non\_chimera\_fasta (fasta) out\_abundance\_biom (biom1) out\_abundance\_count (tabular) 🗇 🤇 summary\_file (html)

#### Chimera

FROGS TSV to BIOM X Abundance TSV File Multi\_hits TSV File biom\_file (biom1) sequence\_file (fasta) 💿 **Convert TSV to** Biom

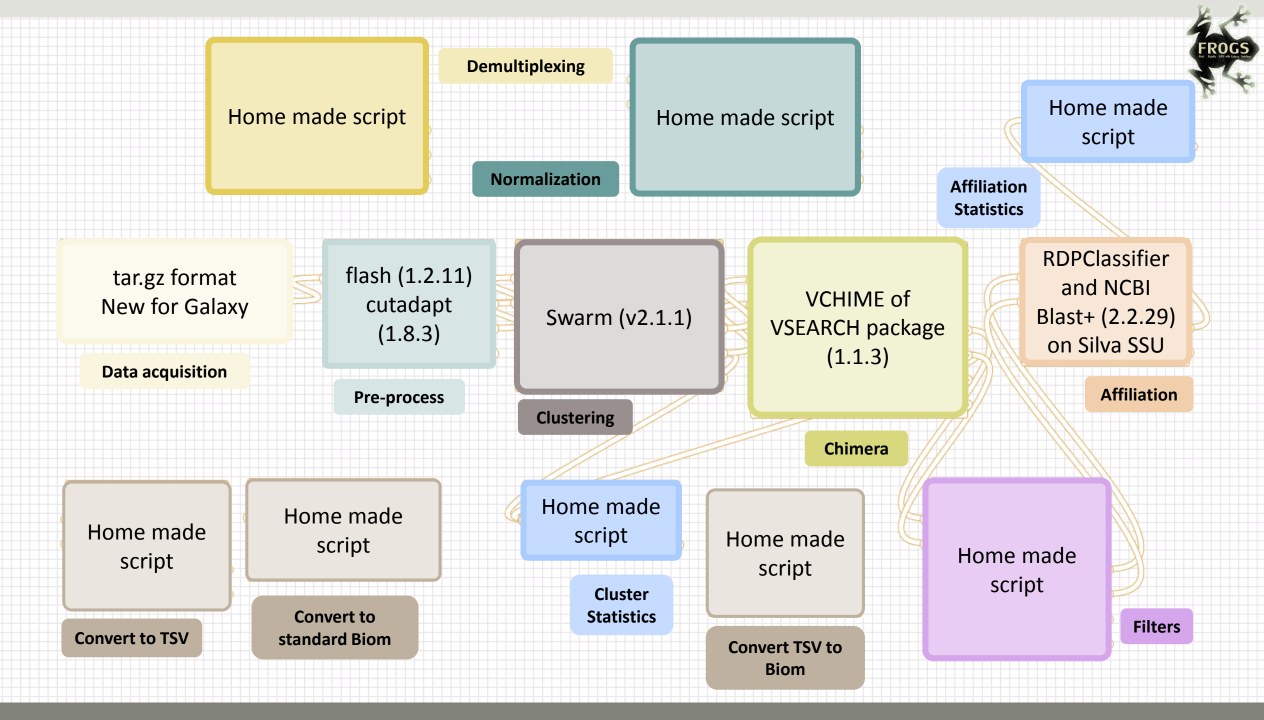
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Affiliation

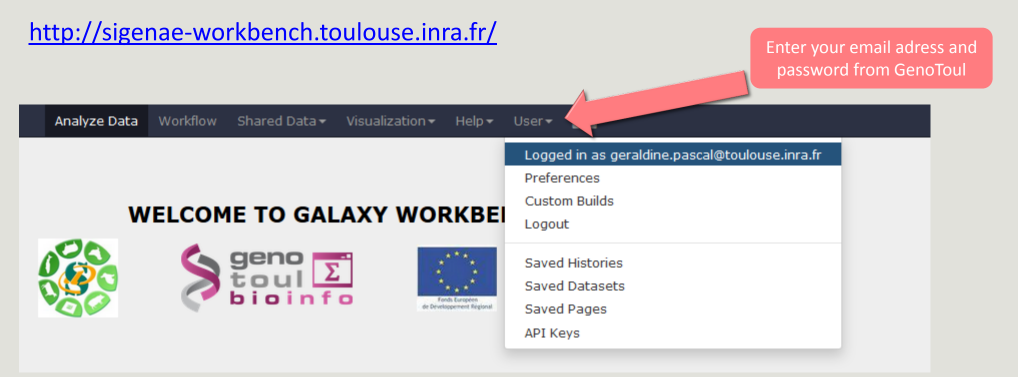
**Filters** 

FROGS Filters Sequences file Abundance file output\_fasta (fasta) output\_biom (biom1) output\_excluded (tabular) ( output\_summary (html)



### Together go to visit FROGS

In your internet browser (Firefox, chrome, Internet explorer) :

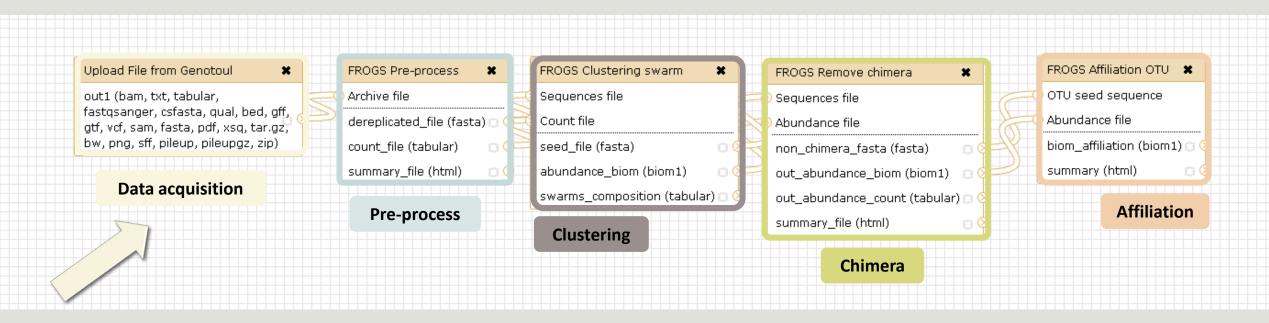


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Tools	FROGS Pre-process Step 1 in metagenomics analysis: denoising and dereplication. (Galaxy Version 1.5.0)	History
METAGENOMICS	Sequencer	search datasets 8
FROGS - Find Rapidly Otu with Galaxy Solution	Illumina	Hantagulumic
FROGS Demultiplex reads	Select the sequencer family used to produce the sequences.	29 shown, 14 <u>deleted</u>
Split by samples the reads in function of inner barcode.	Input type	■ 20.42 MB
FROGS Pre-process Step 1 in	Files by samples	43: FROGS BIOM to std BIOM: ● ✓ ★
metagenomics analysis: denoising and dereplication.	Samples files can be provided in single archive or with two files (R1 and R2) by sample.	blast_metadata.tsv
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FROGS Remove chimera Step 3 in metagenomics analysis :	1: Samples	Abudatasets
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OTU's seed by RDPtools and BLAST	reads 2	38: FROGS
FROGS Clusters stat Process	No fastq dataset available.	Abundance normalisation: report.html
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FROGS Affiliations stat Process some metrics on	+ Insert Samples	Abundance normalisation: normalized.biom
taxonomies.	Reads 1 size	
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FROGS BIOM to TSV Converts	Reads 2 size	30: FROGS BIOM to TSV: multi_hits.tsv
a BIOM file in TSV file.		29: FROGS BIOM to
FROGS TSV to BIOM Converts a TSV file in BIOM file.	The read2 size.	TSV: abundance.tsv
FROGS Abundance	Expected amplicon size	23: FROGS
normalisation		Affiliation OTU:

	<b>=</b> Galaxy	Analyze Data 🛛 Workflow Shared Data 🕶 Visualization 👻 Help 🕶 User 🕶 🗱	Using 5%	
	Tools	FROGS Pre-process Step 1 in metagenomics analysis: denoising and dereplication. (Galaxy Version 1.5.0)   • Options	▲ History 🗶 🌢	
	METAGENOMICS FROGS - Find Rapidly Otu with	Sequencer	FROGS analysis	
	Galaxy Solution	Illumina  Select the sequencer family used to produce the sequences.		
Demultiplexing	FROGS Demultiplex reads Split by samples the reads in function of inner barcode.	Input type	Affiliations stat: summary.html	
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Chimera	3 in metagenomics analysis : Remove PCR chimera in each	1: Samples	<u>②21: FROGS BIOM to</u> ●	
	sample.	Name	©20: FROGS ● Ø X	
Filters	FROGS Filters Filters OTUs on several criteria.	The sample name.	Affiliations stat: summary.html	
A [[]]	FROGS Affiliation OTU Step 4 in metagenomics analysis :	Reads 1	Stat: summary.html	
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Biom to std Biom	taxonomies. FROGS BIOM to std BIOM	Reads 1 size	15: FROGS Filters:	
	Converts a FROGS BIOM in fully compatible BIOM.	The read1 size.	report.html	
Biom to TSV	FROGS BIOM to TSV Converts	Reads 2 size	<u>14: FROGS Filters:</u> ④ Ø 💥 <u>excluded.tsv</u>	
TSV to Biom	a BIOM file in TSV file. FROGS TSV to BIOM Converts	The read2 size.	13: FROGS Filters:	Result files
	a TSV file in BIOM file.	Expected amplicon size	abundance.biom	
Normalization	FROGS Abundance normalisation		<u>12: FROGS Filters:</u>	

# Upload data

Go to demultiplexing tool



### What kind of data ?

### 4 Upload $\rightarrow$ 4 Histories

#### Multiplexed data

Pathobiomes rodents and ticks

multiplex.fastq

barcode.tabular

454 data

Freshwater sediment metagenome

454.fastq.gz

SRA number • SRR443364 MiSeq R1 fastq + R2 fastq

Farm animal feces metagenome

sampleA\_R1.fastq

sampleA\_R2.fastq

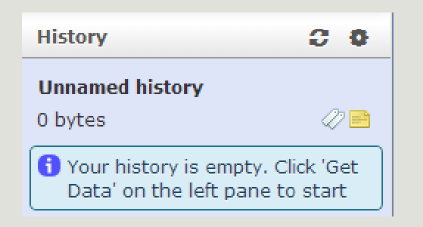
MiSeq contiged fastq in archive tar.gz

Farm animal feces metagenome

100spec\_90000seq\_9s amples.tar.gz

#### 1<sup>ST</sup> CONNEXION

#### **RENAME HISTORY**

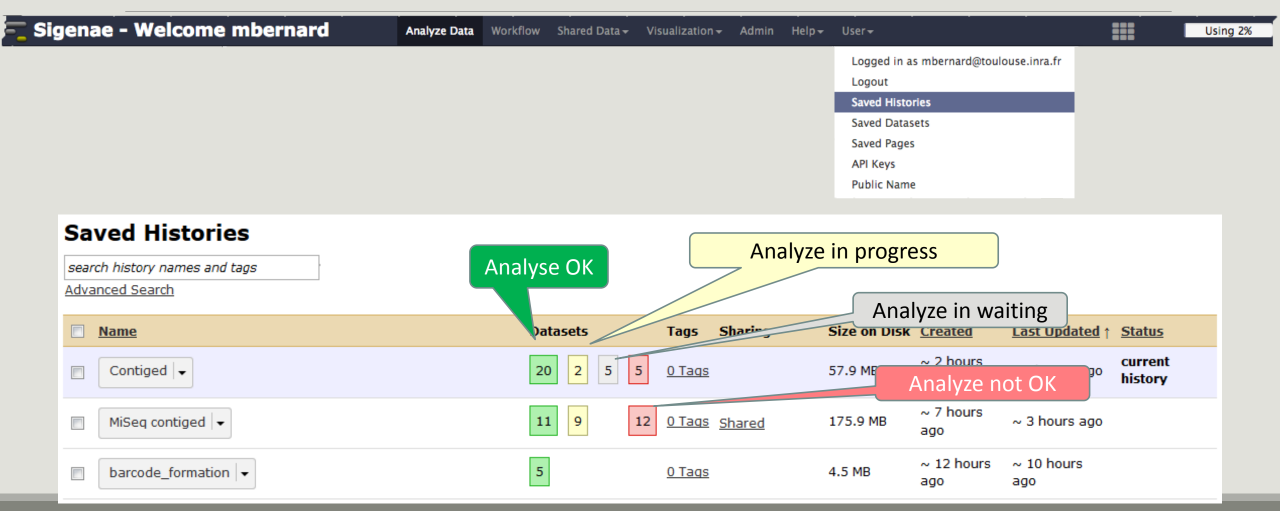


click on Unnamed history, Write your new name, Tap on Enter. 3 0 History Historique renommé 47 🖻 0 bytes 1 Your history is empty. Click 'Get Data' on the left pane to start

### History gestion

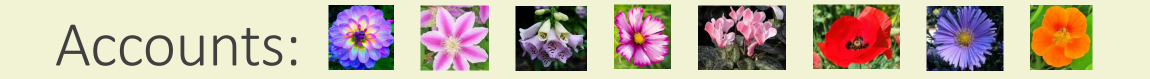
- Keep all steps of your analysis.
- Share your analyzes.
- At each run of a tool, a new dataset is created. The data are not overwritten.
- Repeat, as many times as necessary, an analysis.
- All your logs are automatically saved.
- Your published histories are accessible to all users connected to Galaxy (Shared Data / Published Histories).
- Shared histories are accessible only to a specific user (History / Option / Histories Shared With Me).
- To share or publish a history: User / Saved histories / Click the history name / Share or Publish

Saved Histories



# Your turn! - 1

LAUNCH UPLOAD TOOLS



- anemone
- arome
- aster
- bleuet
- camelia
- capucine
- chardon
- clematite
- cobee

- coquelicot
- cosmos

Password: f1o2r3!

### Your turn: exo 1

Create the 1st history multiplexed

Import files « multiplex.fastq » and « barcode.tabular » present in the Genotoul folder /work/formation/FROGS/

Create the 2nd history 454

Import file « **454.fastq.gz** » present in the **Genotoul** folder /work/formation/FROGS/ (datatype <u>fastq or fastq.gz is the same !)</u>



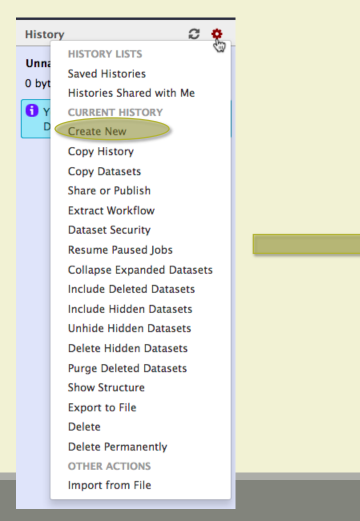
Create the 3rd history MiSeq R1 R2

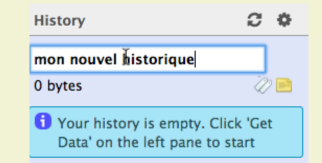
Import files « sampleA\_R1.fastq » and « sampleA\_R2.fastq » present in the Genotoul folder /work/formation/FROGS/

#### Create the 4th history MiSeq contiged

Import archive file « 100spec\_90000seq\_9samples.tar.gz » present in the Genotoul folder /work/formation/FROGS/







Tools	Upload File (version 1.1.3)	
search tools	File Format:	Default method, your files are on
YOUR DATA	Auto-detect	your computer or accessible on the
Upload Data	Which format? See help below	internet, they are copied on your
Upload File	File: Choisissez un fichier Aucun fichier choisi	
Upload File from genotoul	TIP: Due to browser limitations, uploading files larger than 2GB is guaranteed to 1 (below) or FTP (if enabled by the site administrator).	Galaxy account
EBI SRA ENA SRA	URL/Text:	
UCSC Main table browser		
UCSC Test table browser		You can only upload one local file at a time
UCSC Archaea table browser		$\rightarrow$ 10 samples $\geq$ 10 uploads
<u>Get Microbial Data</u>	Here you may specify a list of URLs (one per line) or paste the contents of a file.	You can upload multiple files using URLs
<u>BioMart</u> Central server	Convert spaces to tabs:	
<u>Compress</u> zip or tar file	Yes	but only smaller than 2Go
Download Data	Use this option if you are entering intervals by hand.	



Each uploaded file will consume your Galaxy's quota!

Tools		Upload File (ve	ersion 1.0.0)		
search tools	0	Path to file:			
YOUR DATA			onnees_simulees/100WEPL_setA.tar.gz ike : /work/USERNAME/somewhere/afile		
Upload Data Upload File		File type: tar.gz 🔻	Do not forget to precise the input file type		
Upload File from genotoul		Execute			

Specific SIGENAE GENOTOUL method. It allows you to access to your files in your work account on the Genotoul **without** consuming your Galaxy quota.

And if you have multiple samples ?

See <u>How to create an archiveTAR.ppt</u>



How to transfer files on /work of Genotoul?

See <u>How to transfert to genotoul.ppt</u>

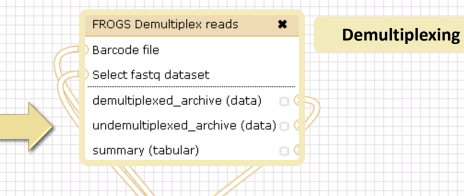
Tools	Upload archive (version 1.0.0)
FROGS - FIND RAPIDLY OTU WITH GALAXY SOLUTION FROGS pipeline Upload archive from your computer Demultiplex reads Split by samples the reads in function of	File: Choisissez un fichier Aucun fichier choisi TIP: Due to browser limitations, uploading files larger than 2GB is guaranteed to fail. To upload large files, use the URL method. URL: Here you may specify the archive URL.
inner barcode. <u>FROGS Pre-process</u> Step 1 in metagenomics analysis (16S/18S): denoising and dereplication.	Execute 3 What it does

If you have an archive on your own computer and smaller than 2Go, you may use this specific FROGS tool to upload your samples archive instead of the default « Upload File » of Galaxy.

Tools	Download from web or upload from disk	
search tools	Regular Composite	
YOUR DATA Upload Data Upload File Upload File from genotoul EBI SRA ENA SRA UCSC Main table browser UCSC Test table browser UCSC Archaea table browser Get Microbial Data	Prop files here	You can only upload multiple files at a time
BioMart Central server	Type (set all): Auto-detect V Q Genome (set all): unspecified (?) V	but only smaller than 2Go
<u>Compress</u> zip or tar file	Chasse legel file Chaste date Deuse Deset Start Class	
Download Data	Choose local file Paste/Fetch data Pause Reset Start Close	

New functionality in latest Galaxy version : <u>http://147.99.108.167/galaxy/</u>

# Demultiplexing tool



#### Upload File from Genotoul × out1 (bam, txt, tabular, fastqsanger, csfasta, qual, bed, gff, gtf, vcf, sam, fasta, pdf, xsq, tar.gz, bw, png, sff, pileup, pileupgz, zip)

### Data acquisition

#### FROGS Pre-process × Archive file dereplicated\_file (fasta) 🖸 count\_file (tabular) seed\_file (fasta) summary\_file (html) **Pre-process**

#### FROGS Clustering swarm Sequences file Count file

abundance\_biom (biom1)

swarms\_composition (tabular) 🖂 🤇

#### Clustering

#### FROGS Remove chimera Sequences file

x

#### Abundance file

×

0(

non\_chimera\_fasta (fasta)

out\_abundance\_biom (biom1) 🛛 🖸 🤅 out\_abundance\_count (tabular) 🗇 🤅

Chimera

summary\_file (html)

#### FROGS Affiliation OTU OTU seed sequence

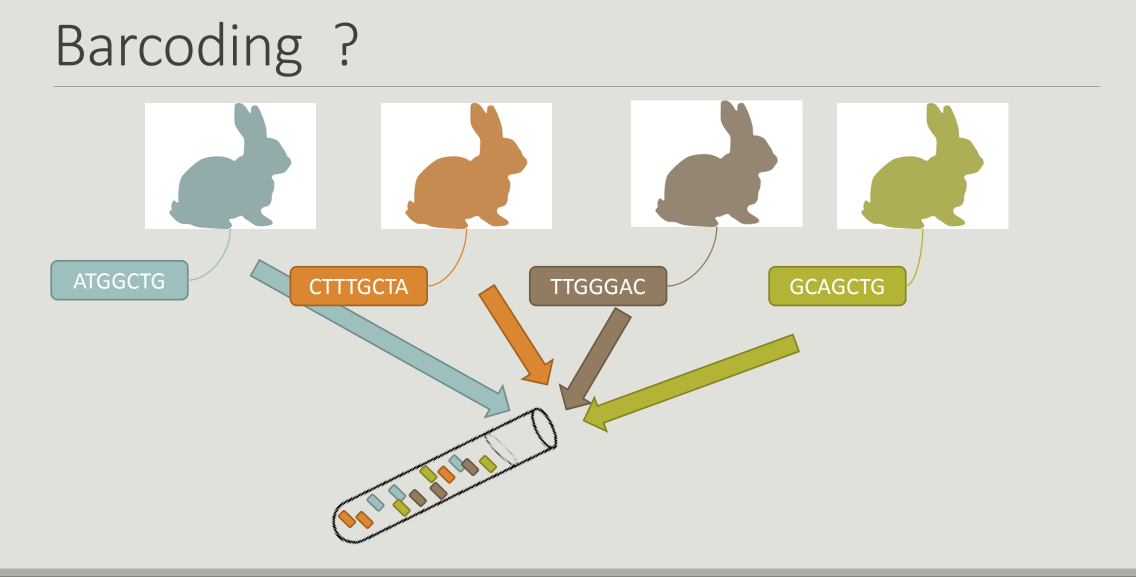
Abundance file

biom\_affiliation (biom1) 🖸

summary (html)

Affiliation

52

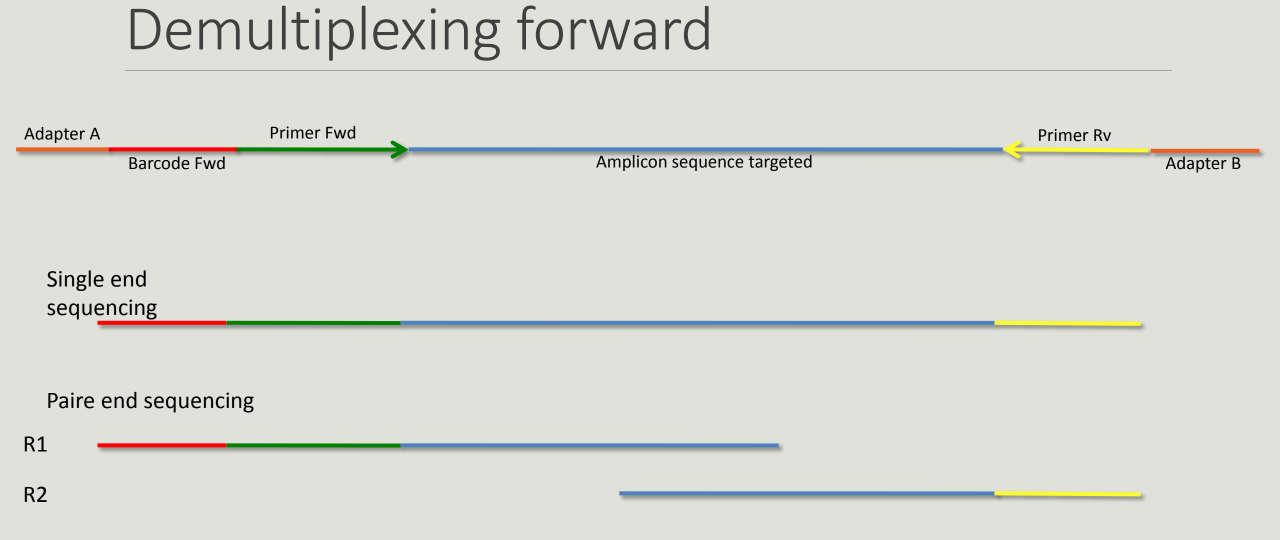


### Demultiplexing

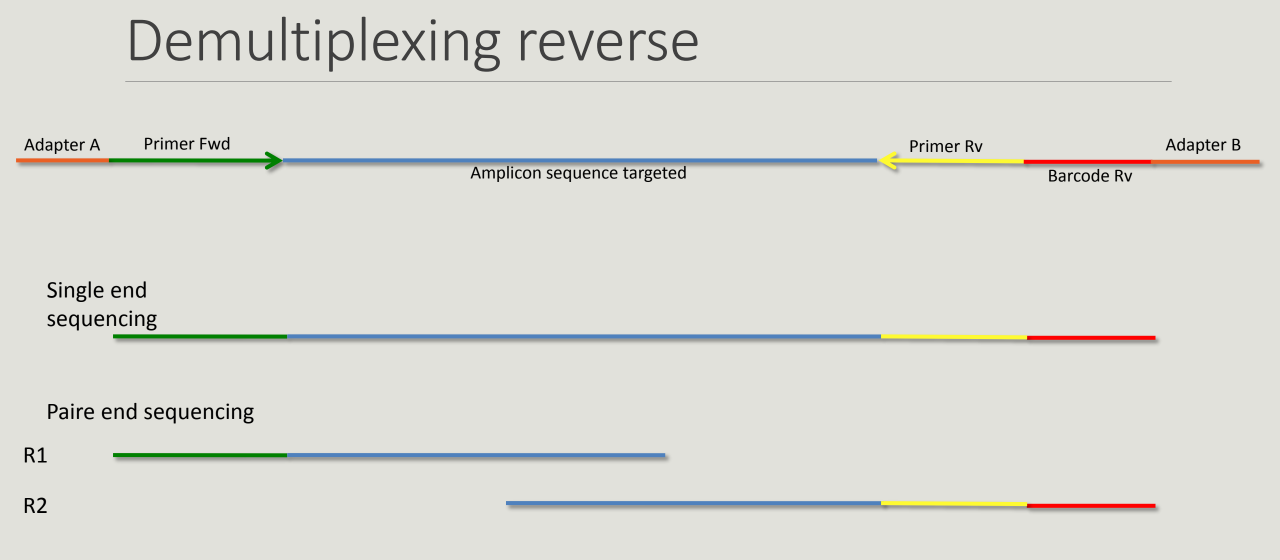
Sequence demultiplexing in function of barcode sequences :

- In forward
- In reverse
- In forward and reverse

Remove unbarcoded or ambiguous sequences

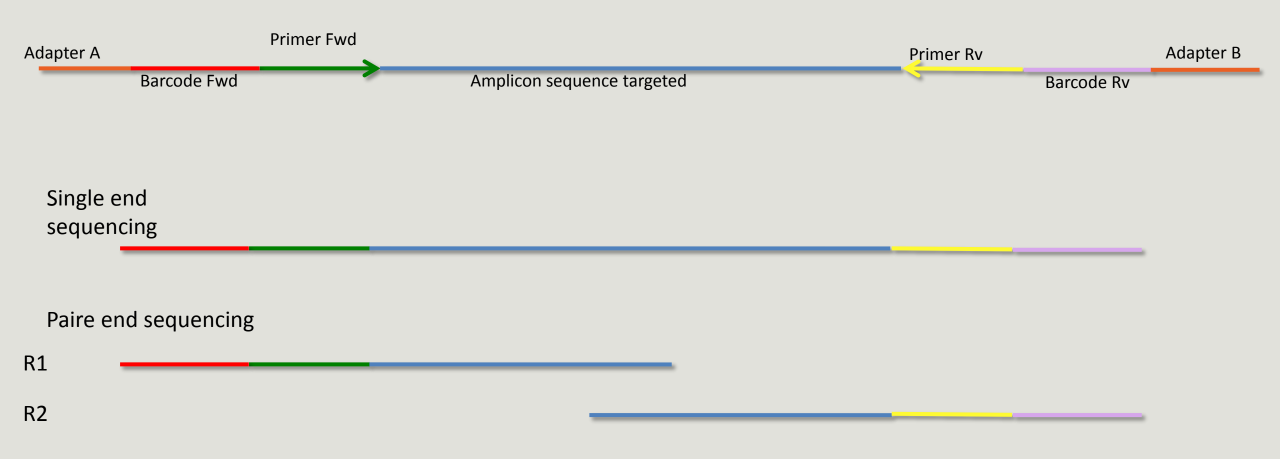


#### 



#### 

### Demultiplexing forward and reverse



# Your turn! - 2

LAUNCH DEMULTIPLEX READS TOOL



- anemone
- arome
- aster
- bleuet
- camelia
- capucine
- chardon
- clematite

- cobee
- coquelicot
- cosmos
- cyclamen

Password: f1o2r3!

#### FROGS Demultiplex reads (version 1.1.0)

#### Barcode file:

#### 1: barcode.tabular 🔻

This file describes barcodes and samples (one line by sample tabulated separated from barcode sequence(s)). See Help section

#### Single or Paired-end reads:

Single 🔻

Select between paired and single end data

#### Select fastq dataset:

Г	,	÷
L	4	_

Specify dataset of your single end reads

#### barcode mismatches:

Number of mismatches allowed in barcode

#### barcode on which end ?:

Forward	•	
Forward		at the begining of the forward end or of the reverse end or both?
Reverse		· ·
Both ends		
Execute		



#### FROGS Demultiplex reads (version 1.1.0)

#### Barcode file:

#### 1: barcode.tabular 🔻

This file describes barcodes and samples (one line by sample tabulated separated from barcode sequence(s)). See Help section

#### Single or Paired-end reads:

Paired 🔻

Select between paired and single end data

#### Select first set of reads:

Specify dataset of your forward reads

#### Select second set of reads:



Specify dataset of your reverse reads

#### barcode mismatches:



Number of mismatches allowed in barcode

#### barcode on which end ?:



### Exercise 2

In **multiplexed** history launch the demultiplex tool:

« The Patho-ID project, rodent and tick's pathobioms study, financed by the metaprogram INRA-MEM, studies zoonoses on rats and ticks from multiple places in the world, the co-infection systems and the interactions between pathogens. In this aim, thay have extracted hundreads of or rats and ticks samples from which they have extracted 16S DNA and sequenced them first time on Roche 454 plateform and in a second time on Illumina Miseq plateform. For this courses, they authorized us to publicly shared some parts of these samples. »

Parasites & Vectors (2015) 8:172 DOI 10.1186/s13071-015-0784-7. Detection of Orientia sp. DNA in rodents from Asia, West Africa and Europe. Jean François Cosson, Maxime Galan, Emilie Bard, Maria Razzauti, Maria Bernard, Serge Morand, Carine Brouat, Ambroise Dalecky, Khalilou Bâ, Nathalie Charbonnel and Muriel Vayssier-Taussat

### Exercise 2

In **multiplexed** history launch the demultiplex tool:

Data are single end reads

 $\rightarrow$  only 1 fastq file

Samples are characterized by an association of two barcodes in forward and reverse strands → multiplexing « both ends »

<u>2: /work/frogs</u> /Formation/multiplex.fas	ھ ta	0	8
<u>1: /work/froqs</u> /Formation/barcode.txt	۲	0	×

### Exercise 2

Demultiplex tool asks for 2 files: one « fastq » and one « tabular »

1. Play with pictograms



- 2. Observe how is built a fastq file.
- 3. Look at the stdout, stderr when available (in the 1) pictogram )

FROGS Demultiplex reads (version 1.1.0)	History 🖸	0
Barcode file:  1: barcode.tabular  This file describes barcodes and samples (one line by sample tabulated separated from	FROGS multiplexed	? 📑
barcode sequence(s)). See Help section   Single or Paired-end reads:   Single    Select between paired and single end data   Select fastq dataset:	2: multiplex.fastq  2.1 MB format: fastqsanger, database: ? Epilog : job finished at Fri Nov 6 15:08:03 CET 2015  (1) (2) (2) (2) (2) (2) (3) (3) (3) (4) (4) (4) (4) (4) (4) (4) (4) (4) (4	/ 🗙
2: multiplex.fastq - Specify dataset of your single end reads	@HNHOSKDØ1ALDØH ATCTAGTGATAAGTTCCGTTCATCCTAAGTCCAT	TATT
barcode mismatches: 0 Number of mismatches allowed in barcode	+ FFFFFFFFFDDA554444889422=<>400044 @HNHOSKD01B8SLE ATAGCTGATTGGTTTAAGCGGATAGGGATTAGAT	
barcode on which end ?: Both ends 💌 The barcode is at the begining of the forward end or of the reverse end or both?	1: barcode.tabular 👁 🕼	
Execute	10 lines format: tabular, database: ? Epilog : job finished at Fri Nov 6 15:07:53 CET 2015	28
🕽 What it does	1 2 3 MgArd0001 ACAGCGT TGTACGT	
Classify single or paired end reads in function of barcode forward or reverse in the first or both reads.	NgArd0009 ACAGTAG TGTACGT MgArd0009 ACGTCAG TGTACGT	
Command line:	MgArd0029 ACTCAGT TGTACGT MgArd0038 ACTCGTC TGTACGT	
demultiplex.pyinput-R1 *FQ_INPUT1* [input-R2 *FQ_INPUT2*]input-barcode *TXT_F	MgArd0046 AGCAGTC TGTACGT	

demultiplex.py --input-R1 \*FQ\_INPUT1\* [--input-R2 \*FQ\_INPUT2\*] --input-barcode \*TXT F

### Advices

#### For your own data

- Do not forget to indicate barcode sequence as they are in the fastq sequence file, especially if you have data multiplexed via the reverse strand.
- For the mismatch threshold, we advised you to let the threshold to 0, and if you are not satisfied by the result, try with 1. The number of mismatch depends on the length of the barcode, but often those sequences are very short so 1 mismatch is already more than the sequencing error rate.
- If you have different barcode lengths, you must demultiplex your data in different times beginning by the longest barcode set and used the "unmatched" or "ambiguous" sequence with smaller barcode and so on.
- If you have Roche 454 sequences in sff format, you must convert them with some program like sff2fastq

### Results

		$\rightarrow$
	A tar archive is created	
	A tar archive is created by grouping one (or a	
	pair of) fastq file per	
	sample with the names indicated in the first	
_	column of the barcode	
	tabular file	

	#sample	count
	ambiguous	0
	MgArd0009	65
	MgArd0017	152
	MgArd0038	1185
	MgArd0029	172
>	unmatched	492
	MgArd0001	85
	MgArd0081	209
	MgArd0046	373
	MgArd0054	217
	MgArd0073	454
	MgArd0062	1109

With barcode mismatches >1 sequence can corresponding to several samples. So these sequences are non-affected to a sample.

Sequences without known barcode. So these sequences are non-affected to a sample.

### Format: Barcode

BARCODE FILE is expected to be tabulated:

- first column corresponds to the sample name (unique, without space)
- second to the forward sequence barcode used (None if only reverse barcode)
- optional third is the reverse sequence barcode (optional)

Take care to indicate sequence barcode in the strand of the read, so you may need to reverse complement the reverse barcode sequence. Barcode sequence must have the same length.

Example of barcode file.

The last column is optional, like this, it describes sample multiplexed by both fragment ends.

MgArd00001 ACAGCGT ACGTACA

### Format : FastQ

FASTQ : Text file describing biological sequence in 4 lines format:

- first line start by "@" correspond to the sequence identifier and optionally the sequence description. "@Sequence\_1 description1"
- second line is the sequence itself. "ACAGC"
- third line is a "+" following by the sequence identifier or not depending on the version
- fourth line is the quality sequence, one code per base. The code depends on the version and the sequencer

@HNHOSKD01ALD0H ACAGCGTCAGAGGGGGTACCAGTCAGCCATGACGTAGCACGTACA + CCCFFFFFFHHHHHJJIJJJJHHFF@DEDDDDDDD@CDDDDACDD

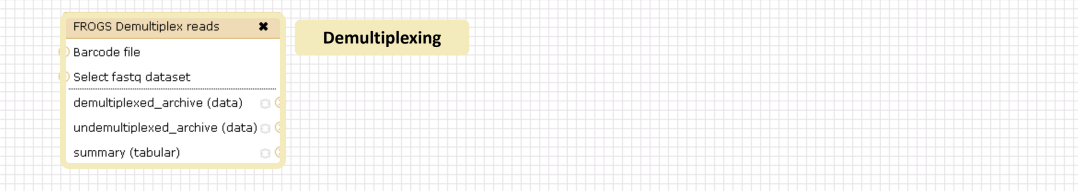
### How it works ?

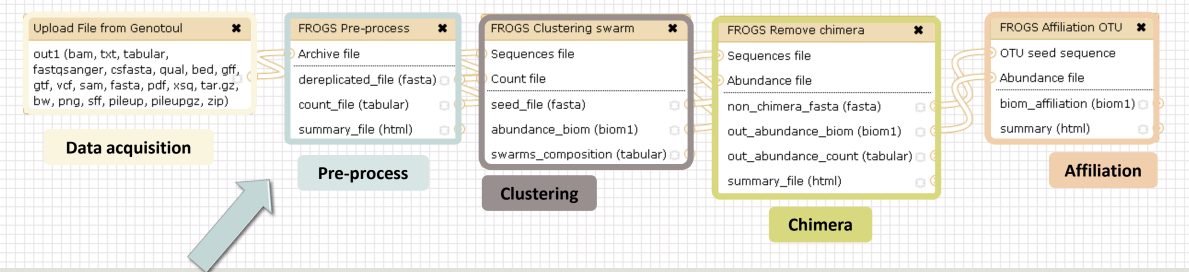
For each sequence or sequence pair the sequence fragment at the beginning (forward multiplexing) of the (first) read or at the end (reverse multiplexing) of the (second) read will be compare to all barcode sequence.

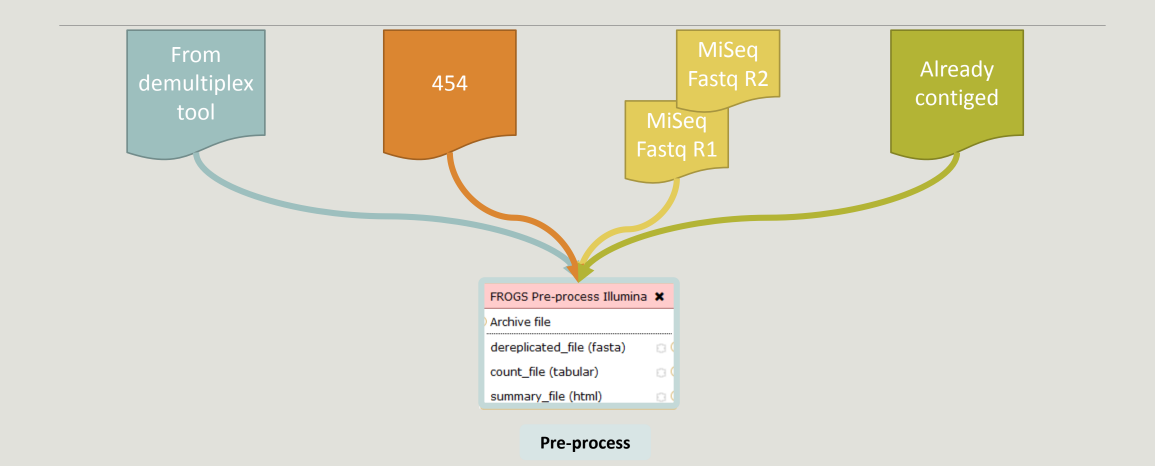
If this fragment is equal (with less or equal mismatch than the threshold) to one (and only one) barcode, the fragment is trimmed and the sequence will be attributed to the corresponding sample.

Finally fastq files (or pair of fastq files) for each sample are included in an archive, and a summary describes how many sequence are attributed for each sample.

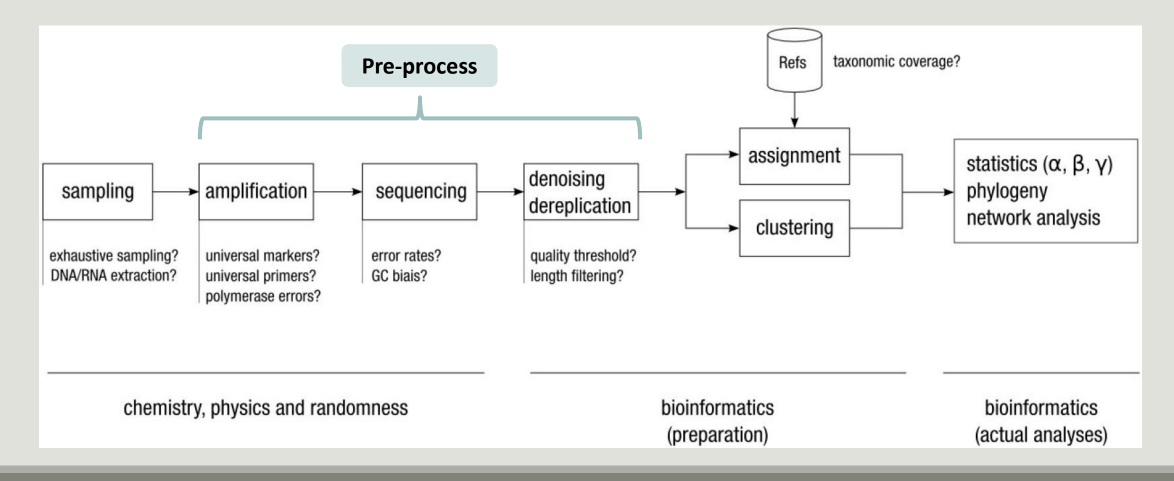
# Pre-process tool







### Amplicon-based studies general pipeline



### Pre-process

- Delete sequence with not expected lengths
- Delete sequences with ambiguous bases (N)
- Delete sequences do not contain good primers
- Dereplication

- + removing homopolymers (size = 8) for 454 data
- + quality filter for 454 data

### Example for:

- Illumina MiSeq data
- 1 sample
- Non joined

Pre-process example 1

OGS Pre-process         Step 1 in metagenomics analysis: denoising and dereplication. (Galaxy Version 1.5.0) <ul></ul>						
quencer						
umina	•					
lect the sequencer family used to produce the sequences.						
input type						
Files by samples	-					
amples files can be provided in single archive or with two files (R1 and R2) by sample.						
Reads already contiged ?						
No	-					
The inputs contain 1 file by sample : Reads 1 and Reads 2 are already contiged by pair.						
Samples						
1: Samples						
Name						
sampleA						
The sample name.						
Reads 1						
1: /work/formation/FROG: /sampleA_R1.fastq	•					
R1 FASTQ file of paired-end reads.						
reads 2						
C 2: /work/formation/FROGS /sampleA_R2.fastq	•					
R2 FASTQ file of paired-end reads.						
+ Insert Samples						
Reads 1 size						
250						
The read1 size.	)					
Reads 2 size						
Parameters for the						
The read2 size.	]					
Expected amplicon size						
410						

Maximum amplicon length expected in approximately 90% of the amplicons.

Minimum amplicon size	
340	
The minimum size for the amplicons.	
Maximum amplicon size	[V3 – V4] 16S variability
450	
The maximum size for the amplicons.	
Sequencing protocol	
Illumina standard	•
The protocol used for sequencing step: standard or	custom with PCR primers as sequencing primers.
5' primer	
CCGTCAATTC	
The 5 primer sequence (wildcards are accepted).	The orienta ameters'. Primer sequences
3' primer	rimer sequences
CCGCNGCTGCT	
The 3' primer sequence (wildcards are accepted).	The orientation is detailed below in 'Primers parameters'.
✓ Execute	

### Example for:

- Sanger 454 data
- 1 sample
- Joined

### Pre-process example 2

ROGS Pre-process Step 1 in metagenomics analysis: denoising and dereplication. (Galaxy Version 1.5.0)	<ul> <li>Options</li> </ul>
equencer	
54	•
elect the sequencer family used to produce the sequences.	
Input type	
One file by sample	•
Samples files can be provided in single archive or with one file by sample.	
Samples	
1: Samples	
Name	
my_sample	
The sample name.	
Sequence file	
🗋 🙆 🗀 1: /work/formation/FROGS/454.fastq.gz	
FASTQ file of sample.	
+ Insert Samples	
Minimum amplicon size	
380	
The minimum size for the amplicons (with primers).	
Maximum amplicon size [V3 – V4] 16S variability	
500	
The maximum size for the amplicons (with primers).	
5' primer	
ACGGGAGGCAGCAG	
The 5 primer sequence (wildcards are accepted). The orient ameters'.	
3' primer Primer sequences	
AGGATTAGATACCCTGGTA	
The 3' primer sequence (wildcards are accepted). The orientation is detailed below in Primers parameters'.	

### **Pre-process example 3**

Execute

Archive file

Sequencer

Input type

Archive

Illumina

• Without sequenced PCR primers (Kozich protocol)

The tar file containing the sequences file(s	;) for each sample.
Yes	Paire-end sequencing all ready joined
The archive contains 1 file by sample : Rea	ads 1 and Reads 2 are already contiged by pair.
linimum amplicon size	
380	
he minimum size for the amplicons.	[V3 – V4] 16S variability
laximum amplicon size	
500	
he maximum size for the amplicons.	
equencing protocol	
Custom protocol (Kozich et al. 2013)	No more primers
he protocol used for sequencing step: stan	dard or custom with PCR primers as sequencing primers.

**Sequencing technology** 

One file per sample and all files are contained in a archive

### Example for:

Joined

- Illumina MiSeq data
- 9 samples in 1 archive

Samples files can be provided in single archive or with two files (R1 and R2) by sample.

Select the sequencer family used to produce the sequences.

•

-

# Your turn! - 3

GO TO EXERCISES 3

Go to « 454 » history

Launch the pre-process tool on that data set

 $\rightarrow$  objective : understand the parameters

1- Test different parameters for « minimum and maximum amplicon size »

2- Enter these primers: Forward: ACGGGAGGCAGCAG Reverse: AGGATTAGATACCCTGGTA

#### FROGS Pre-process (version 1.4.2)

#### Sequencer:

```
454
```

Select the sequencer family used to produce the sequences.

### Input type:

One file by sample 🔻

Samples files can be provided in single archive or with one file by sample.

#### Samples

Samples 1	Sample name is require					
Name:						
my_sample						
The sample name.						

#### Sequence file:

6: /work/formation/FROGS/454.fastq.gz ▼ FASTQ file of sample.

### Add new Samples

#### Minimum amplicon size:

380

The minimum size for the amplicons (with primers).

### Maximum amplicon size:

### 500

The maximum size for the amplicons (with primers).

### 5' primer:

ACGGGAGGCAGCAG

The 5' primer sequence (wildcards are accepted). The orientation is detailed

### 3' primer:

Execute

### AGGATTAGATACCCTGGTA

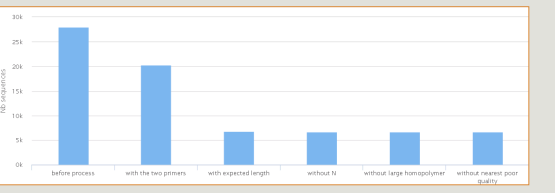
The 3' primer sequence (wildcards are accepted). The orientation is detailed

Primers used for sequencing V3-V4: Forward: ACGGGAGGCAGCAG Reverse: AGGATTAGATACCCTGGTA

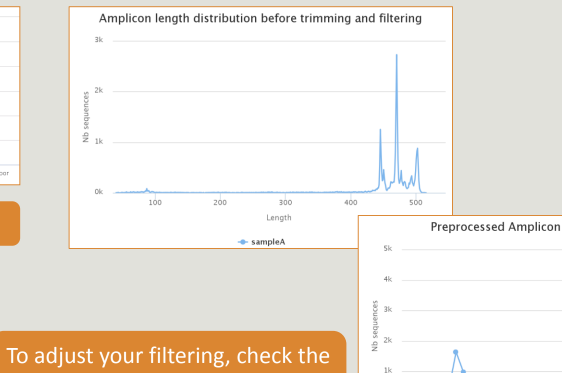
### Size range of 16S V3-V4: [ 380 - 500 ]

What do you understand about amplicon size, which file can help you ?
What is the length of your reads before preprocessing ?
Do you understand how enter your primers ?
What is the « FROGS Pre-process: dereplicated.fasta » file ?
What is the « FROGS Pre-process: count.tsv » file ?
Explore the file « FROGS Pre-process: report.html »
Who loose a lot of sequences ?

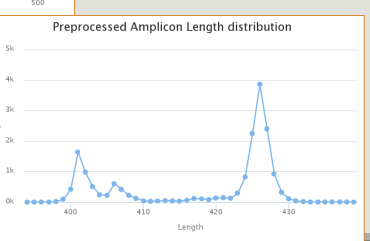
	Samples	before process ∲	with the two primers	with expected length	without N	without large ¢	without nearest poor quality
	sample_454	28,009	20,227	6,806	6,677	6,675	6,672



To be kept, sequences must have the 2 primers

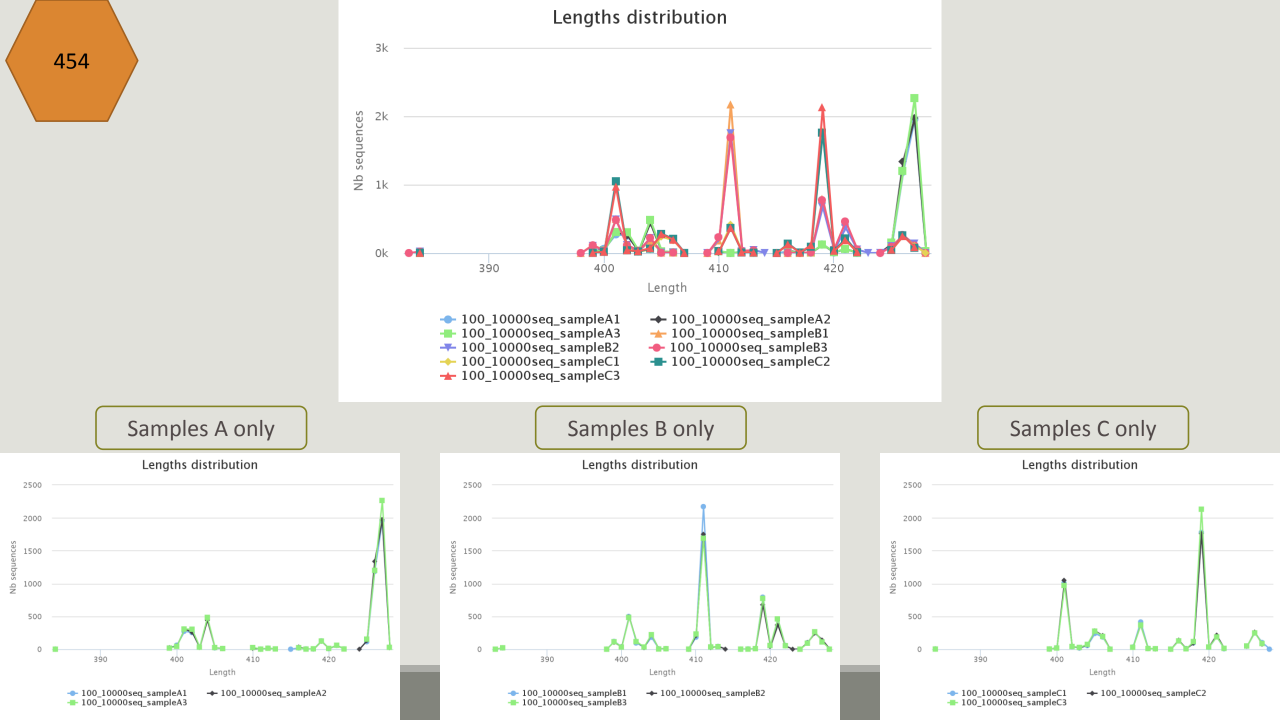


distribution of sequence lengths.



--- sampleA

454



## Cleaning, how it work ?

Filter contig sequence on its length which must be between min-amplicon-size and maxamplicon-size

use cutadapt to search and trim primers sequences with less than 10% differences

Minimum am	plico	n si	ize:		
380					
The minimum	size	for	the	amplico	ons.
Maximum am	plico	on s	ize:		

500

\_ \_ \_ \_

The maximum size for the amplicons.

## Cleaning, how it work ?

dereplicate sequences and return one uniq fasta file for all sample and a count table to indicate sequence abundances among sample.

In the HTML report file, you will find for each filter the number of sequences passing it, and a table that details these filters for each sample.



Go to « MiSeq R1 R2 » history

- Launch the pre-process tool on that data set
- $\rightarrow$  objective: understand flash software

#### FROGS Pre-process (version 1.4.2)

#### Sequencer:

### Illumina 🝷

Select the sequencer family used to produce the sequences.

### Input type:

### Files by samples 💌

Samples files can be provided in single archive or with two files (R1 and R2) by sample.

### Reads already contiged ?:

```
No 💌
```

The inputs contain 1 file by sample : Reads 1 and Reads 2 are already contiged by pair.

### Samples

### Samples 1

### Name:

sampleA

The sample name.

### Reads 1:

1: /work/formation/FROGS/sampleA\_R1.fastq 💌

R1 FASTQ file of paired-end reads.

### reads 2:

2: /work/formation/FROGS/sampleA\_R2.fastq 💌

R2 FASTQ file of paired-end reads.

### Add new Samples

### Reads 1 size:

250 The read1 size.

### Reads 2 size:

250 The read2 size.

### Primers used for this sequencing : Forward: CCGTCAATTC Reverse: CCGCNGCTGCT

Lecture 5'  $\rightarrow$  3'

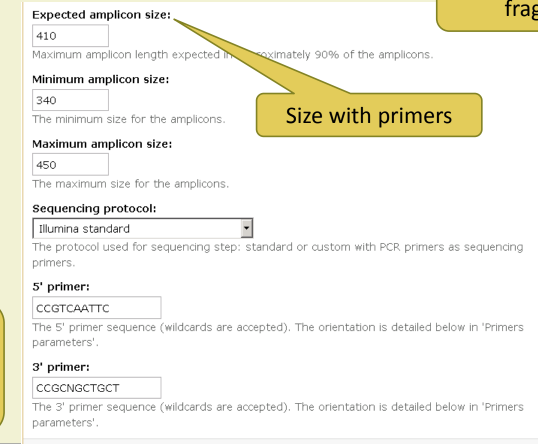
### >ERR619083.M00704

CGCTTGCCACCTACGTATTACCGCNGCTGCT

### Real 16S sequenced fragment

MiSeq

R1 R2



Execut

## Flash, how it works ?

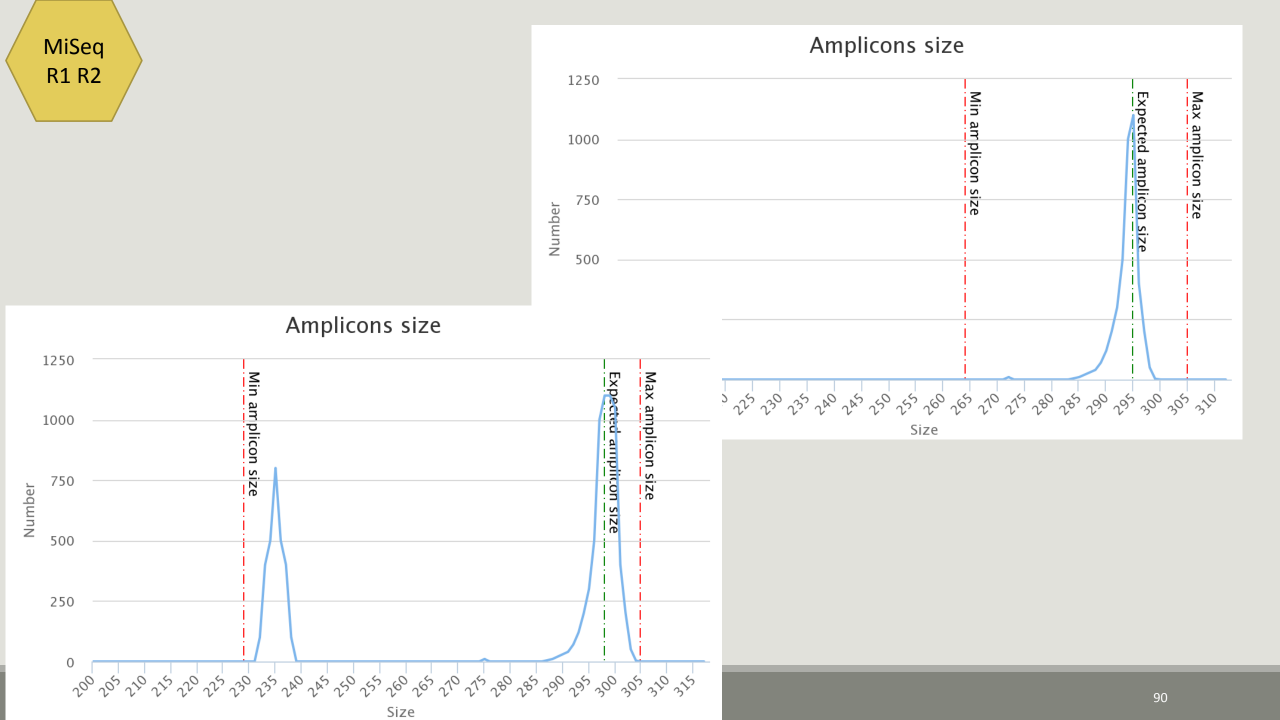
To contig read1 and read2 with FLASh with :

- a minimum overlap equals to
- [(R1-size + R2-size) expected-amplicon-size]

ex: minimum overlap (250+250) - 410 = 50 maximum overlap 450

and a maximum overlap equal to [expected-amplicon-size] with a maximum of 10% mismatch among this overlap

90% of the amplicon are smaller than [expected-amplicon-size]





Interpret « FROGS Pre-process: report.html » file.



Go to« MiSeq contiged » history

- Launch the pre-process tool on that data set
- $\rightarrow$  objective: understand output files



3 samples are **technically replicated** 3 times : 9 samples of 10 000 sequences each.

100\_10000seq\_sampleA1.fastq100\_10000seq\_sampleB1.fastq100\_10000seq\_sampleC1.fastq100\_10000seq\_sampleA2.fastq100\_10000seq\_sampleB2.fastq100\_10000seq\_sampleC2.fastq100\_10000seq\_sampleA3.fastq100\_10000seq\_sampleB3.fastq100\_10000seq\_sampleC3.fastq



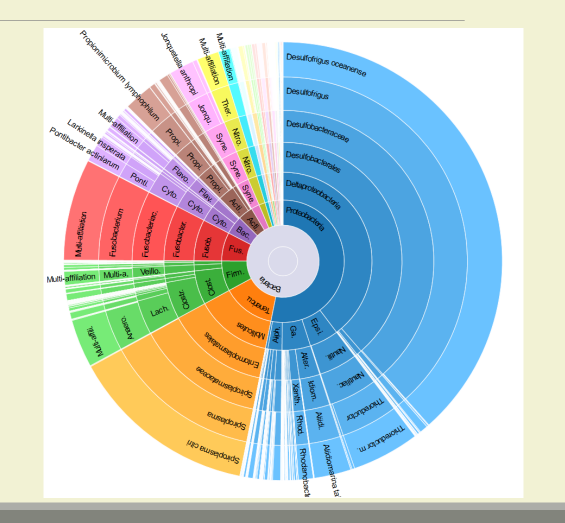
- 100 species, covering all bacterial phyla
- Power Law distribution of the species abundances
- Error rate calibrated with real sequencing runs
- 10% chimeras
- 9 samples of 10 000 sequences each (90 000 sequences)

Normal

Distribution

Power Law

Distribution





"Grinder (v 0.5.3) (Angly et al., 2012) was used to simulate the PCR amplification of full-length (V3-V4) sequences from reference databases. The reference database of size 100 were generated from the LTP SSU bank (version 115) (Yarza et al., 2008) by

- (1) filtering out sequences with a N,
- (2) keeping only type species
- (3) with a match for the forward (ACGGRAGGCAGCAG) and reverse (TACCAGGGTATCTAATCCTA) primers in the V3-V4 region and
- (4) maximizing the phylogenetic diversity (PD) for a given database size. The PD was computed from the NJ tree distributed with the LTP."

### FROGS Pre-process (version 1.4.2)

#### CO Sequencer:

IV

#### Illumina 👻

Select the sequencer family used to produce the sequences.

#### Input type:

#### Archive

Samples files can be provided in single archive or with two files (R1 and R2) by sample.

#### Archive file:

1: /work/formation/FROGS/100spec\_90000seq\_9samples.tar.gz 
The tar file containing the sequences file(s) for each sample.

#### Reads already contiged ?:

### Yes 🔻

The archive contains 1 file by sample : Reads 1 and Reads 2 are already contiged by pair.

#### Minimum amplicon size:

#### 380

The minimum size for the amplicons.

#### Maximum amplicon size:

#### 500

The maximum size for the amplicons.

#### Sequencing protocol:

Illumina standard

The protocol used for sequencing step: standard or custom with PCR primers as sequencing primers.

#### 5' primer:

#### ACGGGAGGCAGCAG

The 5' primer sequence (wildcards are accepted). The original

### 3' primer:

### TAGGATTAGATACCCTGGT

The 3' primer sequence (wildcards are accepted). The ori

Primers used for this sequencing : 5' primer: ACGGGAGGCAGCAG 3' primer: TAGGATTAGATACCCTGGTA Lecture 5'  $\rightarrow$  3'

#### Lengths distribution $\equiv$ ₽ 1k 385 390 395 405 410 415 420 425 Length 100\_10000seg\_sampleA1 + 100\_10000seq\_sampleA2 - 100\_10000seq\_sampleA3 100\_10000seq\_sampleB1 100\_10000seq\_sampleB3 100\_10000seq\_sampleC1 = 100\_10000seq\_sampleC2 + 100\_10000seq\_sampleC3

Click on legend

Amplicons lengths

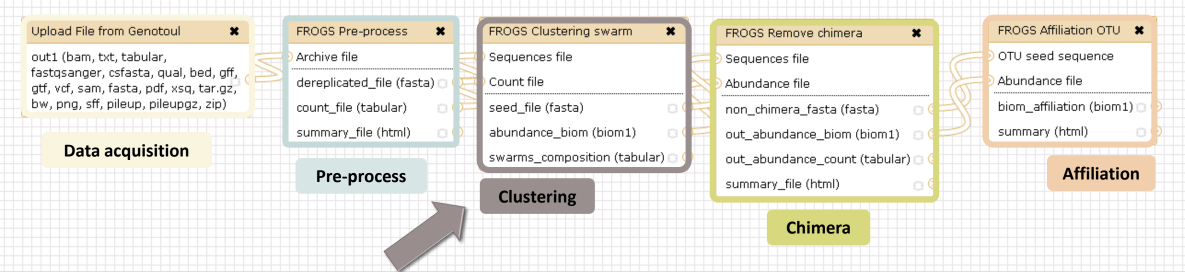
Execute

### Exercise 3.3 - Questions

- 1. How many sequences are there in the input file ?
- 2. How many sequences did not have the 5' primer?
- 3. How many sequences still are after pre-processing the data?
- 4. How much time did it take to pre-process the data ?
- 5. What can you tell about the sample based on sequence length distributions ?

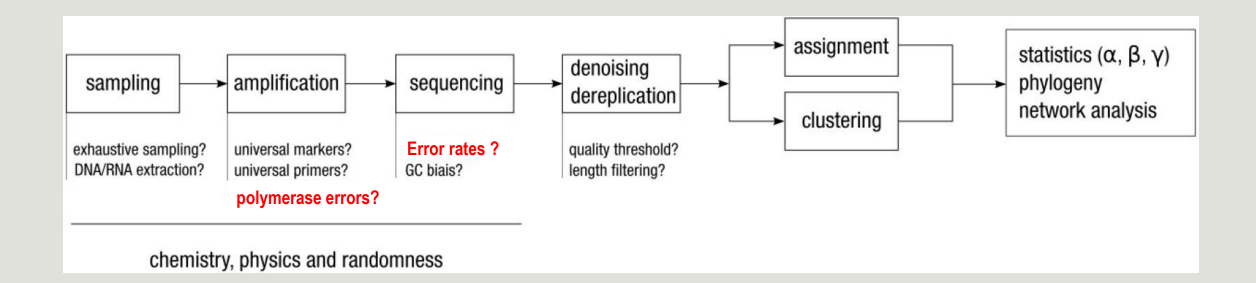
# Clustering tool

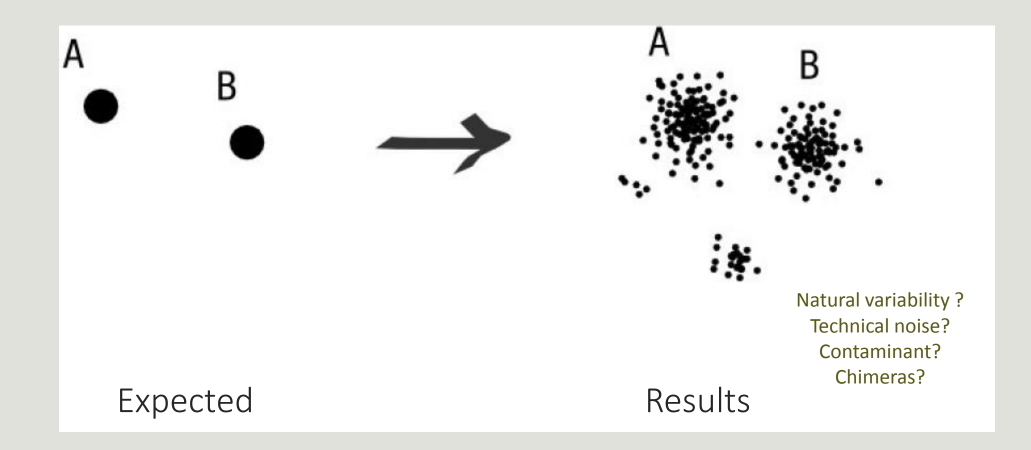


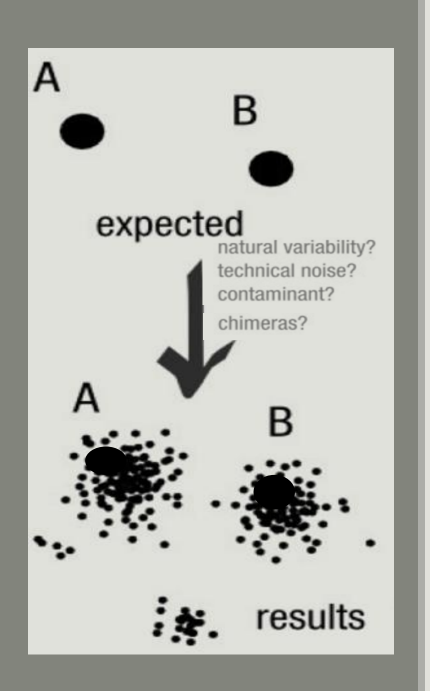


## Why do we need clustering ?

Amplication and sequencing and are not perfect processes







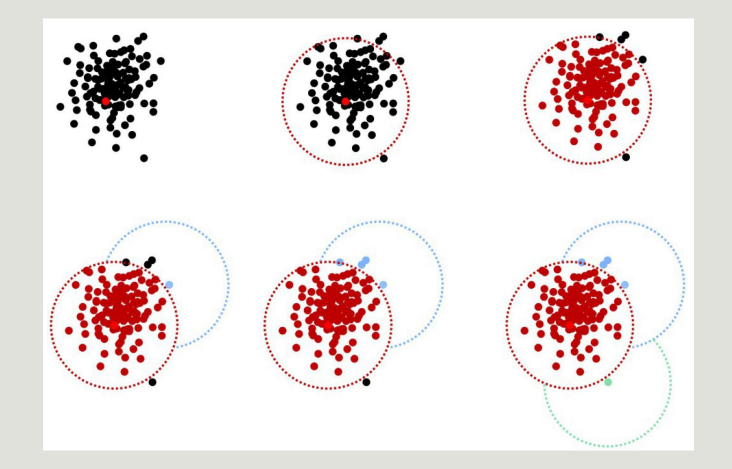
### To have the best accuracy:

### Method: All against all

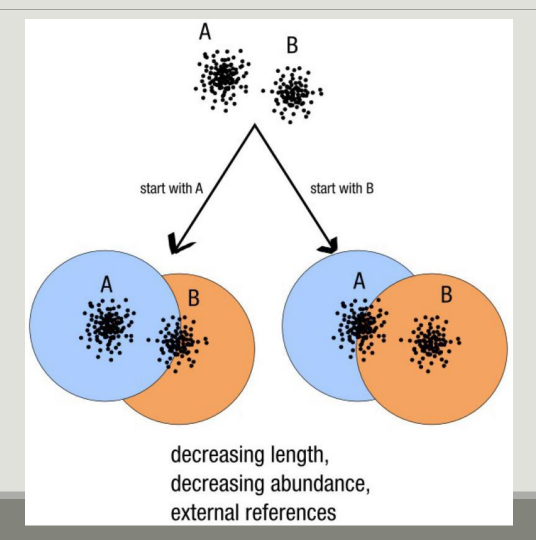
- Very accurate
- Requires a lot of memory and/or time

=> Impossible on very large datasets without strong filtering or sampling

### How traditional clustering works ?

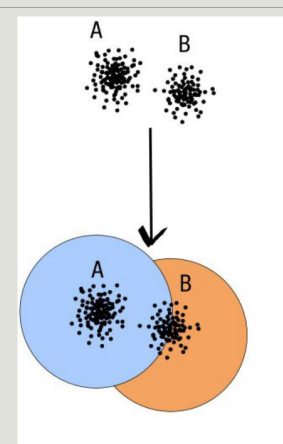


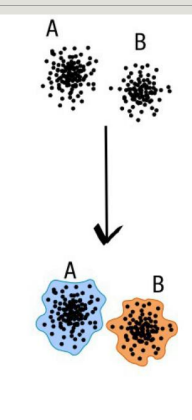
### Input order dependent results



### Fréderic Mahé communication

## Single a priori clustering threshold

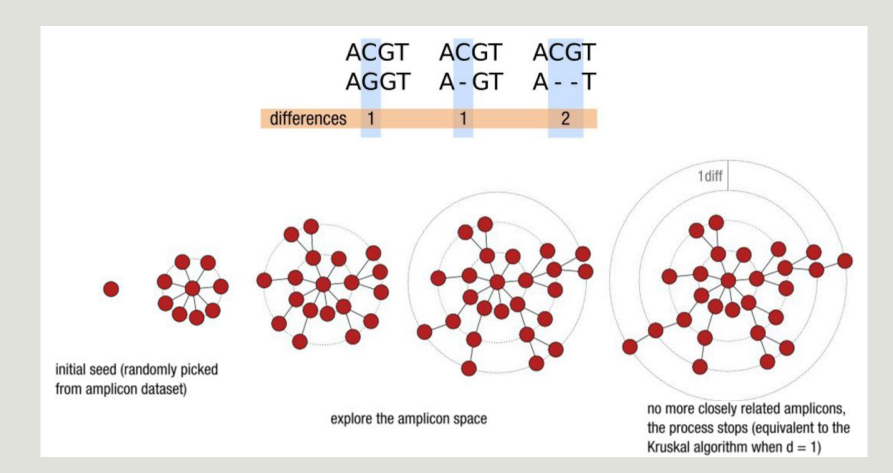




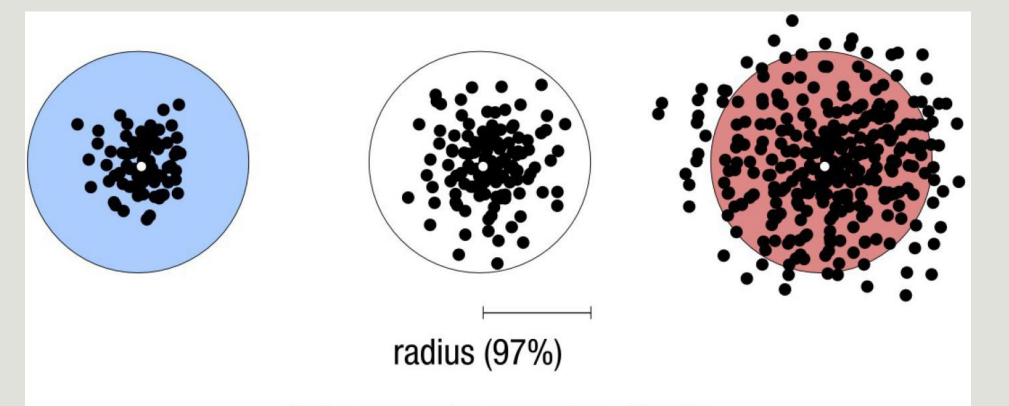
compromise threshold unadapted threshold natural limits of clusters

Fréderic Mahé communication

### Swarm clustering method

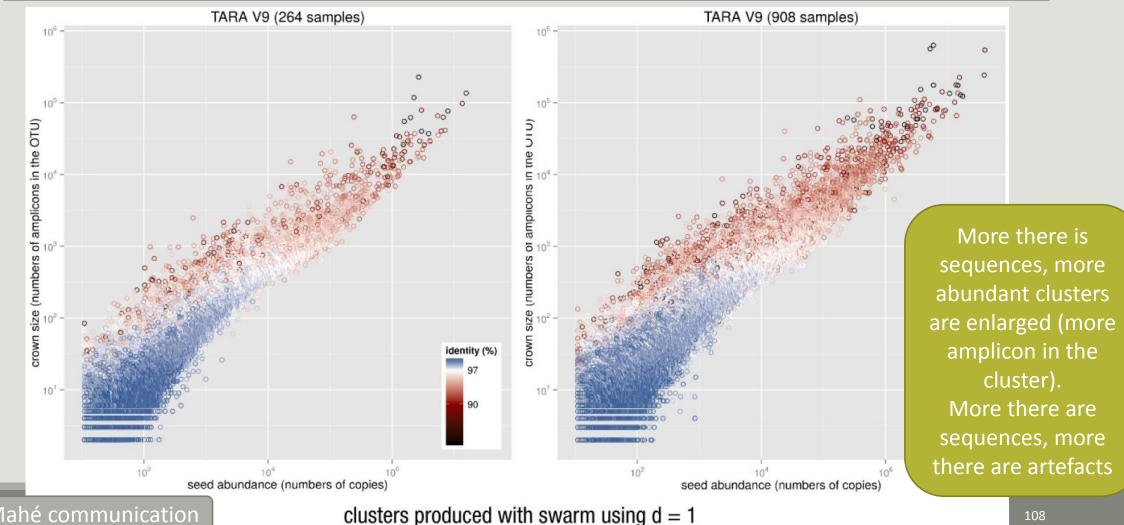


### Comparison Swarm and 3% clusterings



Radius expressed as a percentage of identity with the central amplicon (97% is by far the most widely used clustering threshold)

### Comparison Swarm and 3% clusterings



Fréderic Mahé communication



A robust and fast clustering method for amplicon-based studies.

The purpose of **swarm** is to provide a novel clustering algorithm to handle large sets of amplicons.

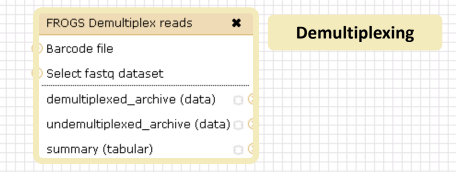
**swarm** results are resilient to input-order changes and rely on a small **local** linking threshold *d*, the maximum number of differences between two amplicons.

swarm forms stable high-resolution clusters, with a high yield of biological information.

Swarm: robust and fast clustering method for amplicon-based studies. Mahé F, Rognes T, Quince C, de Vargas C, Dunthorn M. PeerJ. 2014 Sep 25;2:e593. doi: 10.7717/peerj.593. eCollection 2014. PMID:25276506

FROGS Clustering swarm	FROGS Clustering swarm Step 2 in metagenomics analysis : clustering. (Galaxy Version 2.3.0)	▼ Options		
Sequences file	Sequences file			
Count file	2: FROGS Pre-process: dereplicated.fasta	•		
abundance_biom (txt)	The sequences file (format: fasta).			
seed_file (fasta)	Count file			
swarms_composition (tabular) 🕥	3: FROGS Pre-process: count.tsv	•		
	It contains the count by sample for each sequence (format: TSV).			
Clustering	Aggregation distance			
	3			
	Maximum number of differences between sequences in each aggregation step.			
	Performe denoising clustering step?			
	Yes No If checked, clustering will be perform in two steps, first with distance = 1 and then with your input distance			
	✓ Execute			
	1st run for denoising:	1st run for denoising:		
	Swarm with d = 1 -> high clusters definition			
	linear complexity			
	<u>2<sup>nd</sup> run for clustering:</u> Swarm with d = 3 on the seeds of first Swarm			
	quadratic complexity			
	Gain time !			
	Remove false positives !			

# Cluster stat tool



### Upload File from Genotoul

out1 (bam, txt, tabular, fastqsanger, csfasta, qual, bed, gff, gtf, vcf, sam, fasta, pdf, xsq, tar.gz, bw, png, sff, pileup, pileupgz, zip)

×

### Data acquisition

FROGS Pre-process 🛛 🗙	FROGS Clustering swarm	×	FROGS Remove chimera	K
) Archive file	Sequences file	) Sequences file 📃 🛁		e
dereplicated_file (fasta) 🔅 (	Count file	🔾 Count file		X
count_file (tabular) 🛛 🔅	seed_file (fasta)		non_chimera_fasta (fasta)	
summary_file (html) 🛛 🖸	o abundance_biom (biom1)	0(5)	out_abundance_biom (biom1)	a 🖊
	swarms_composition (tabular	swarms_composition (tabular) 🗅 🤇 🚝		a (
Pre-process	Clustering	Clustering		a 🛛 🔛
			Chimera	
	FROGS Clusters stat 🗶			
	Abundance file			
	summary_file (html) 🛛 🔿			
	Cluster			
	Statistics			

# FROGS Affiliation OTU OTU seed sequence Abundance file biom\_affiliation (biom1) summary (html) Affiliation

✓ Options
•

# Your Turn! - 4

LAUNCH CLUSTERING AND CLUSTERSTAT TOOLS



Go to « MiSeq contiged » history

Launch the Clustering SWARM tool on that data set with aggregation distance = 3 and the denoising

- $\rightarrow$  objectives :
  - understand the denoising efficiency
  - understand the ClusterStat utility



- 1. How much time does it take to finish?
- 2. How many clusters do you get ?



3. Edit the biom and fasta output dataset by adding d1d3

<u>Attributes</u>	Convert Format	<u>Datatype</u>	Permissions
Edit Attribut	es		
Name: warm: seed Info:	d_sequencesd1d3.fa	asta	
/src/galaxy	/usr/local/bioinfo -test/galaxy-	• •	
Annotation	/ Notes:		

FROGS Clusters stat Process some metrics on clusters.

Ø

4. Launch FROGS Cluster Stat tools on the previous abundance biom file

## MiSeq contiged

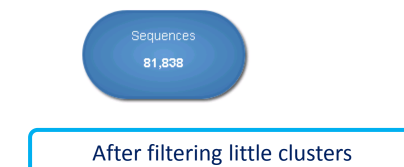
# Exercise 4

- 5. Interpret the boxplot: Clusters size summary
- 6. Interpret the table: **Clusters size details**
- 7. What can we say by observing the **sequence distribution**?
- 8. How many clusters share "sampleB3" with at least one other sample?
- 9. How many clusters could we expect to be shared ?
- **10**. How many sequences represent the 550 specific clusters of "sampleC2"?
- **11**. This represents what proportion of "sampleC2"?
- **12**. What do you think about it?
- **13**. How do you interpret the « Hierarchical clustering » ?

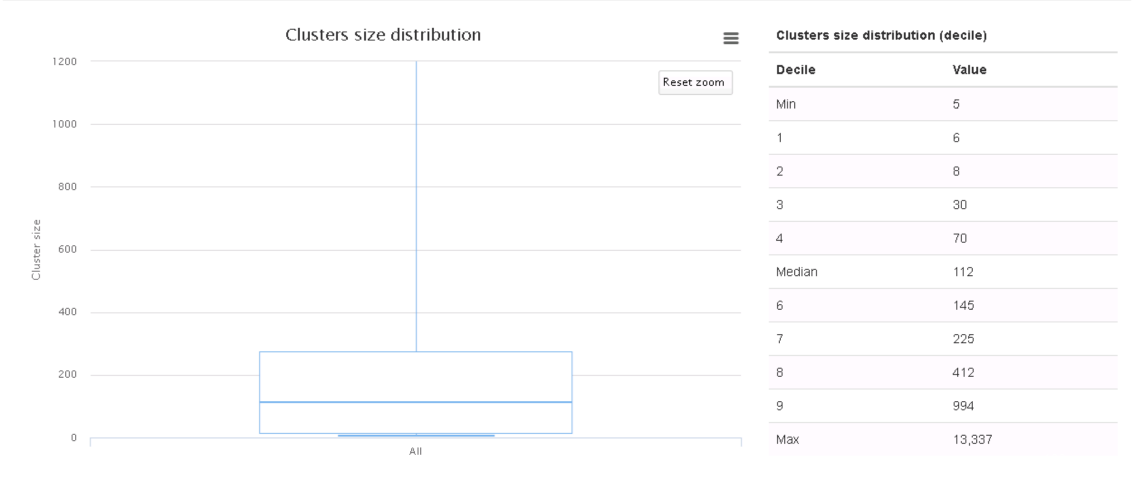
The « Hierachical clustering » is established with a Bray Curtis distance particularly well adapted to abundance table of very heterogenous values (very big and very small figures).

Sigenae - Welcome ı	mbernard	Analyze Data Workflow Shared Data → V	Visualization <del>-</del> Admin F	Help∓ User∓		Using
Tools	Clusters distribution	Sequences distribution Samples distribution				History
deepTools 🔺						15: FROGS Filters: sequences.fasta
FROGS - FIND RAPIDLY OTU WITH GALAXY SOLUTION FROGS pipeline		Clusters	Sequ	uences		14: FROGS Remove
<u>FROGS Upload archive</u> from your computer		5,945	89.	9,721		<u>13: FROGS Remove</u>
FROGS Demultiplex reads Split by samples the reads in function of inner barcode.	/		Mo	st of cluste	ers are singletons	non chimera abundance.bi <u>12: FROGS Remove</u> (2) chimera: non chimera.fasta
<u>FROGS Pre-process</u> Step 1 in metagenomics analysis: denoising and dereplication.	Clusters ?	size summary				11: FROGS Clusters
<u>FROGS Clustering swarm</u> Step 2 in metagenomics analysis : clustering.	15k	Clusters size distribution	≡	Clusters size dist		summary swarm d1d3.htm 102.0 NB format: html, database: <u>?</u>
<u>FROGS Remove chimera</u> Step 3				Decile	Value	## Application Software :/usr/local/bioinfo/src/galaxy-
in metagenomics analysis : Remove PCR chimera in each	12.5k			Min	1	<pre>dev/galaxy-dist/tools/FROGS/t     /clusters_stat.py (version : 1.)</pre>
sample.	12.5K			1	1	Command : /usr/local/bioinfo /src/galaxy-dev/galaxy-dist/to
<u>FROGS Filters</u> Filters OTUs on several criteria.	10k			2	1	/FROGS/tools/clusters_stat.py input-biom /galaxydata
FROGS Affiliation OTU Step 4 in metagenomics analysis :				з	1	/database/file 🔒 🕕 🖓
Taxonomic affiliation of each	Size			4	1	HTML file
OTU's seed by RDPtools and BLAST	at 7.5k			Median	1	
FROGS BIOM to TSV Converts	Ū				•	10: FROGS Clustering
a BIOM file in TSV file.	5k			6	1	<u>swarm:</u> swarms composition d1d3.
FROGS Clusters stat Process some metrics on clusters.				7	1	9: FROGS Clustering
FROGS Affiliations stat Process some metrics on taxonomies.	2.5k			8	2	swarm: abundance_d1d3.bi
FROGS BIOM to std BIOM				9	2	8: FROGS Clustering  Swarm:
Converts a FROGS BIOM in fully compatible BIOM.	0k			Max	13,337	seed sequences d1d3.fasta
FROGS Abundance normalisation		All				7: FROGS Pre-process:
<b></b>						- FROMO D
						110

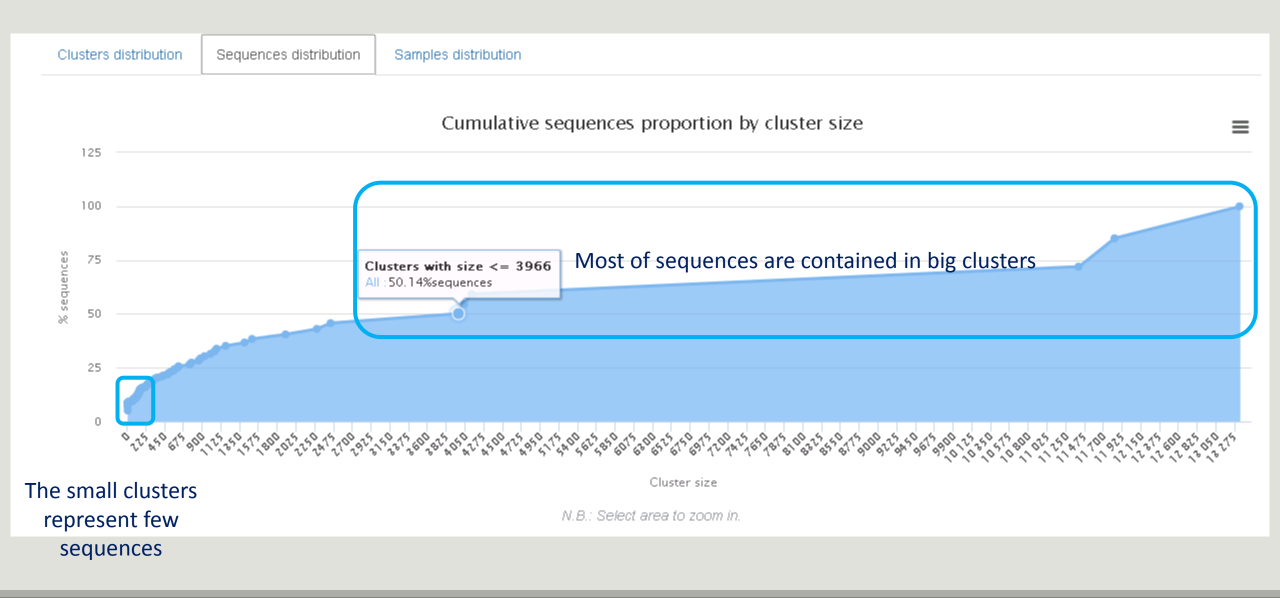




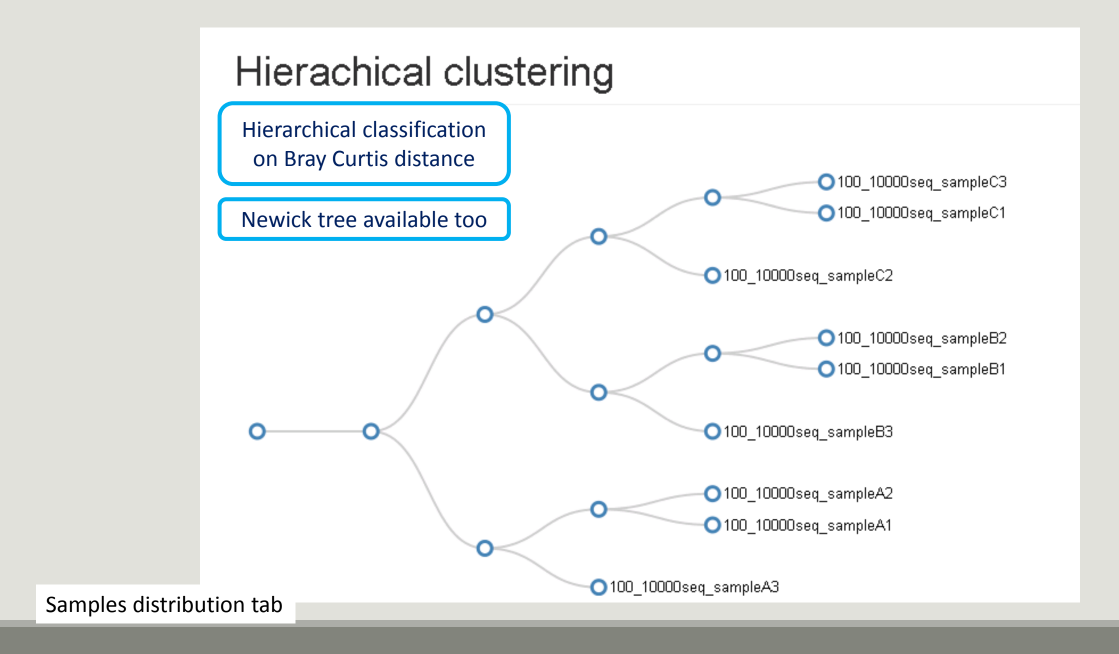
# Clusters size summary



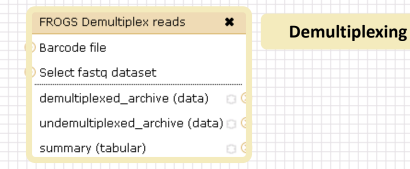
Clusters size details			
		Most of clusters are singletons	
Show 10 - entries		Search:	
Clusters size Cluster size	Number of cluster	♦ % of all clusters	
1	4,595	77.36	
2	866	14.58	
3	155	2.61	
4 After	83	1.40	
5 clustering	42	0.71	
6	29	0.49	
7	22	0.37	
8	13	0.22	
9	6	0.10	
10	6	0.10	



Sequences	- 367 clusters of sampleA1	58 % of the specific clusters of sampleA1				
Show 10    entries Samples information	are common at least once with another sample	represent around 5% of sequences Could be interesting to remove if individual variability is not the concern of user			kcsv	
Sample	Shared clusters	Own clusters	Shared sequences	Own sequences	÷	
100_10000seq_sampleA1	367	513	9,447	528		
100_10000seq_sampleA2	365	490	9,476	503		
100_10000seq_sampleA3	384	483	9,478	494		
100_10000seq_sampleB1	395	548	9,397	572		
100_10000seq_sampleB2	375	508	9,455	515		
100_10000seq_sampleB3	376	562	9,388	579		
100_10000seq_sampleC1	372	539	9,413	552		
100_10000seq_sampleC2	389	550	9,408	567		
100_10000seq_sampleC3	361	516	9,442	525		
Showing 1 to 9 of 9 entries				Previous 1	Next	



# Chimera removal tool



### Upload File from Genotoul

out1 (bam, txt, tabular, fastqsanger, csfasta, qual, bed, gff, gtf, vcf, sam, fasta, pdf, xsq, tar.gz, bw, png, sff, pileup, pileupgz, zip)

×

### **Data acquisition**

FROGS Pre-process 🛛 🗙	FROGS Clustering swarm	FROGS Remove chimera
Archive file	Sequences file	Sequences file
dereplicated_file (fasta) 🖸	Count file	Abundance file
count_file (tabular) 🛛 🖸	🔎 seed_file (fasta) 🛛 🛛 🖓	non_chimera_fasta (fasta)
summary_file (html) 🛛 🖸	🔎 abundance_biom (biom1) 👘 🛛 🕤	out_abundance_biom (biom
	swarms_composition (tabular)	out_abundance_count (tabu
Pre-process	Clustering	summary_file (html)
	FROGS Clusters stat <b>¥</b> Abundance file	Chimera
	summary_file (html)	
	Cluster Statistics	Su Su

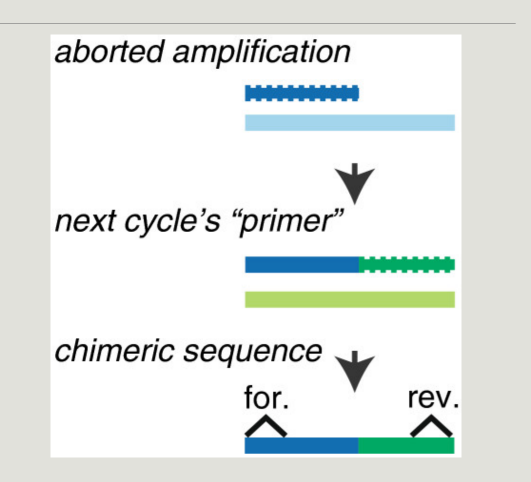
### FROGS Affiliation OTU × OTU seed sequence Abundance file biom\_affiliation (biom1) summary (html) nce\_biom (biom1) ince\_count (tabular) 🖂 🤇 Affiliation

Our advice: **Removing Chimera after** Swarm denoising + Swarm d=3, for saving time without sensitivity loss

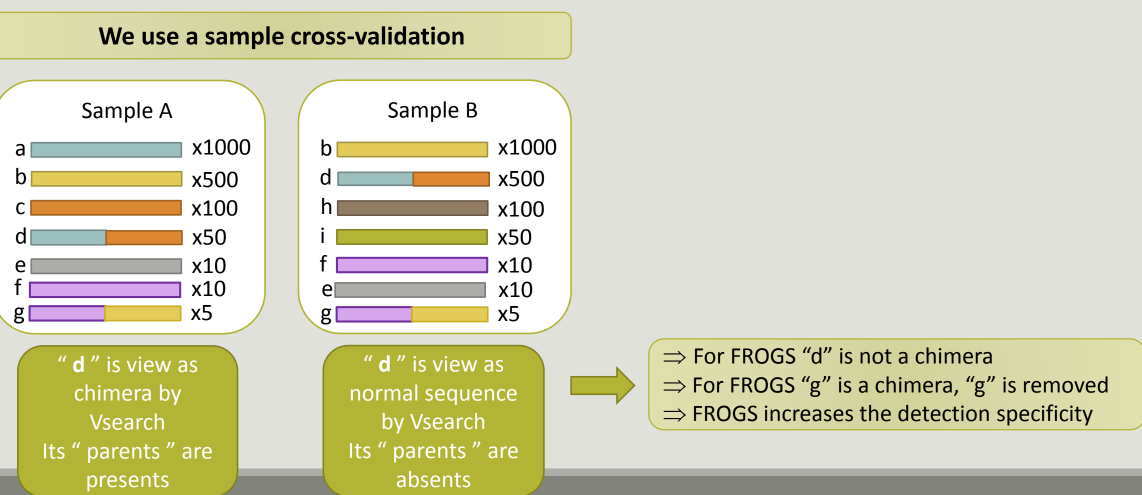
# What is chimera ?

PCR-generated chimeras are typically created when an aborted amplicon acts as a primer for a heterologous template. Subsequent chimeras are about the same length as the non-chimeric amplicon and contain the forward (for.) and reverse (rev.) primer sequence at each end of the amplicon.

**Chimera: from 5 to 45% of reads** (Schloss 2011)



# A smart removal chimera to be accurate



# Your Turn! - 5

LAUNCH THE REMOVE CHIMERA TOOL



Go to « MiSeq contiged » history

Launch the « FROGS Remove Chimera » tool

Follow by the « FROGS ClusterStat » tool on the swarm d1d3 non chimera abundance biom

 $\rightarrow$  objectives :

- understand the efficiency of the chimera removal
- make links between small abundant OTUs and chimeras

	FROGS Remove chimera Step 3 in metagenomics analysis : Remove PCR chimera in each sample. (Galaxy Version 1.3.0)	tions
FROGS Remove chimera 🗶	Sequences file         Image: Step Step Step Step Step Step Step Step	•
non_chimera_fasta (fasta) 💿 🤅	Abundance type BIOM file	•
out_abundance_biom (biom1) 💿 🤇 out_abundance_count (tabular) 💼 🤇	Select the type of file where the abundance of each sequence by sample is stored. Abundance file	
summary_file (html) 💿 🤅	Aduitance me         Image: Construction of the second se	•
Chimera	It contains the count by sample for each sequence.	



- 1. Understand the « FROGS remove chimera : report.html»
  - a. How many clusters are kept after chimera removal?
  - b. How many sequences that represent ? So what abundance?
  - c. What do you conclude ?

## MiSeq contiged

# Exercise 5

- 2. Launch « FROGS ClusterStat » tool on non\_chimera\_abundanced1d3.biom
- 3. Rename output in summary\_nonchimera\_d1d3.html
- 4. Compare the HTML files
  - a. Of what are mainly composed singleton ? (compare with precedent summary.html)
  - b. What are their abundance?
  - c. What do you conclude ?

The weakly abundant OTUs are mainly false positives, our data would be much more exact if we remove them

# Filters tool



### Upload File from Genotoul × out1 (bam, txt, tabular, fastqsanger, csfasta, qual, bed, gff,

gtf, vcf, sam, fasta, pdf, xsq, tar.gz, bw, png, sff, pileup, pileupgz, zip)

### Data acquisition

FROGS Pre-process 🗶 🗔	FROG
Archive file	) Seque
dereplicated_file (fasta) 💿 🤤	Count
count_file (tabular) 🛛 💿 🖙	seed_
summary_file (html) 🛛 💿 🗘 🗖	abund
	swarn
Pre-process	Clu

Demultiplexing

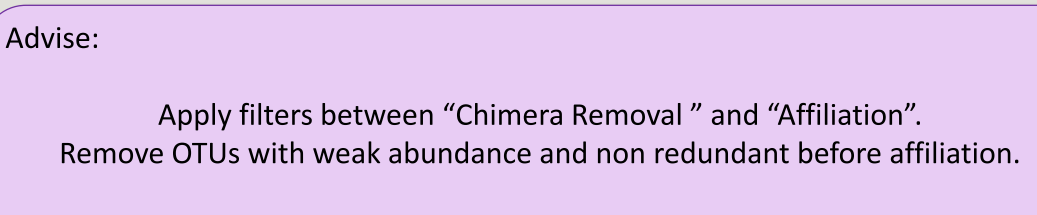
#### S Clustering swarm × FROGS Remove chimera × ences file Sequences file it file Abundance file \_file (fasta) non\_chimera\_fasta (fasta) 00 dance biom (biom1) 0( out abundance biom (biom1) 🛛 🔅 ms\_composition (tabular) | out\_abundance\_count (tabular) 🗇 🤇 summary\_file (html) stering Chimera FROGS Clusters stat X Abundance file summary\_file (html) 📋 Cluster **Statistics** output\_excluded (tabular) 🗇

## FROGS Affiliation OTU OTU seed sequence Abundance file biom\_affiliation (biom1) 🖂 summary (html) Affiliation FROGS Filters × Sequences file Abundance file output\_fasta (fasta) 0( output\_biom (biom1)

output\_summary (html)

**Filters** 

Affiliation runs long time



You will gain time !

# Filters

Filters allows to filter the result thanks to different criteria et may be used after different steps of pipeline :

- On the abundance
- On RDP affiliationOn Blast affiliation
- On phix contaminant

FROGS Filters	×
Sequences file	
Abundance file	
output_fasta (fasta)	Û
output_biom (biom1)	8
output_excluded (tabular)	8
output_summary (html)	8

## Filters

# 4 filter sections

FROGS Filters Filters OTUs on several criteria. (Galaxy Version 1.2.0)	✓ Options
Sequences file	
C & C	•
9: FROGS Remove chimera: non_chimera.fasta	
The sequence file to filter (format: fasta).	
Abundance file	
10: FROGS Remove chimera: non_chimera_abundance.biom	
The abundance file to filter (format: BIOM).	
*** THE FILTERS ON OTUS IN SAMPLES, OTUS SIZE and SEQUENCE PERCENTAGE	
	A la una da na a filta na
Apply filters	Abundance filters
If you want to filter OTUs on their abundance and occurrence.	
Minimum number of samples	
Fill the field only if you want this treatment. Keep OTU present in at least this number of samples.	
Minimum proportion/number of sequences to keep OTU	
Fill the field only if you want this treatment. Use decimal notation for proportion (example: 0.01 for keep OTU with at least 19	
Use integer notation for number of sequence (example: 2 for keep OTU with at least 2 sequences, so remove single singleton	).
N biggest OTU	
Fill the fields only if you want this treatment. Keep the N biggest OTU.	
*** THE FILTERS ON RDP	
	RDP affiliation filters
Apply filters If you want to filter OTUs on their taxonomic affiliation produced by RDP.	NUP anniation miters
Rank with the bootstrap filter	
Nothing selected	<b>-</b>
Minimum bootstrap % (between 0 and 1)	
*** THE FILTERS ON BLAST	
Apply filters	BLAST affiliation filters
If you want to filter OTUs on their taxonomic affiliation produced by Blast.	
Maximum e-value (between 0 and 1)	
Fill the field only if you want this treatment	
Minimum identity % (between 0 and 1)	
Fill the field only if you want this treatment	
Minimum coverage % (between 0 and 1)	
Fill the field only if you want this treatment	
Minimum alignment length	
Fill the field only if you want this treatment	
*** THE FILTERS ON CONTAMINATIONS	
Apply filters	Contamination filter
If you want to filter OTUs on classical contaminations.	
Cotaminant databank	
phiX	
prix The phiX databank (the phiX is a control added in Illumina sequencing technologies).	
f huns anneans faire huns is a countral against an submitting sedaction d countral dept.	
✓ Execute	

# Input

FROGS Filters Filters OTUs on several criteria. (Galaxy Version 1.2.0)	✓ Options		
Sequences file			
9: FROGS Remove chimera: non_chimera.fasta			
The sequence file to filter (format: fasta).	Fasta sequences and its		
Abundance file	corresponding abundance biom files		
10: FROGS Remove chimera: non_chimera_abundance.b	iom		
The abundance file to filter (format: BIOM).			

# Filter 1 : abundance

** THE FILTERS ON OTUS IN SAMPLES, OTUS SIZE and SEQUENCE PERCENTAGE
Apply filters
you want to filter OTUs on their abundance and occurrence.
Minimum number of samples
3
Fill the field only if you want this treatment. Keep OTU present in at least this number of samples.
Minimum proportion/number of sequences to keep OTU
0.00005
Fill the field only if you want this treatment. Use decimal notation for proportion (example: 0.01 for keep OTU with at least 1% of all sequences) ; Use integer notation for number of sequence (example: 2 for keep OTU with at least 2 sequences, so remove single singleton).
N biggest OTU
100
Fill the fields only if you want this treatment. Keep the N biggest OTU.

*** THE FILTERS ON RDP				
Apply filters	<b>•</b>			
you want to filter OTUs on their taxonomic affiliation produced by RDP.				
Rank with the bootstrap filter	Filter 2 & 3:			
Genus				
Minimum bootstrap % (between 0 and 1)	affiliation			
0.8	)			
*** THE FILTERS ON BLAST				
Apply filters	•			
you want to filter OTUs on their taxonomic affiliation produced by Blast.				
Maximum e-value (between 0 and 1)				
Fill the field only if you want this treatment				
Minimum identity % (between 0 and 1)				
1				
Fill the field only if you want this treatment				
Minimum coverage % (between 0 and 1)				
0.95	)			
Fill the field only if you want this treatment				
Minimum alignment length				
Fill the field only if you want this treatment				



	Cotaminant databank
	phiX
	The phiX databank (the phiX is a control added in Illumina sequencing technologies).

Soon, several contaminant banks

# Your Turn! - 6

LAUNCH DE LA TOOL FILTERS



Go to history « MiSeq contiged »

Launch « Filters » tool with non\_chimera\_abundanced1d3.biom, non\_chimerad1d3.fasta Apply 2 filters :

- Minimum proportion/number of sequences to keep OTU: 0.00005\*
- Minimum number of samples: 3

 $\rightarrow$  objective : play with filters, understand their impacts on falses-positives OTUs

#### FROGS Filters

### Sequences file

Abundance file output\_fasta (fasta)

output\_biom (biom1)

×

8

output\_excluded (tabular) 🖂 🤇

output\_summary (html) 🛛 🔅

### Filters

### Apply filters

If you want to filter OTUs on their abundance and occurrence.

FROGS Filters Filters OTUs on several criteria. (Galaxy Version 1.2.0)

🗋 🔁 🗀 10: FROGS Remove chimera: non\_chimera\_abundance.biom

\*\*\* THE FILTERS ON OTUS IN SAMPLES, OTUS SIZE and SEQUENCE PERCENTAGE

🕒 🙆 🗅 9: FROGS Remove chimera: non\_chimera.fasta

#### Minimum number of samples

The sequence file to filter (format: fasta).

The abundance file to filter (format: BIOM).

#### 3

Sequences file

Abundance file

Fill the field only if you want this treatment. Keep OTU present in at least this number of samples.

#### Minimum proportion/number of sequences to keep OTU

#### 0.00005

Fill the field only if you want this treatment. Use decimal notation for proportion (example: 0.01 for keep OTU with at least 1% of all sequences); Use integer notation for number of sequence (example: 2 for keep OTU with at least 2 sequences, so remove single singleton).

#### N biggest OTU

Fill the fields only if you want this treatment. Keep the N biggest OTU.

#### \*\*\* THE FILTERS ON RDP

#### No filters

If you want to filter OTUs on their taxonomic affiliation produced by RDP.

#### \*\*\* THE FILTERS ON BLAST

No filters

If you want to filter OTUs on their taxonomic affiliation produced by Blast.

#### \*\*\* THE FILTERS ON CONTAMINATIONS

No filters

If you want to filter OTUs on classical contaminations.

🗸 Execute

### Output

92: FROGS Filters: report.html	• / %
91: FROGS Filters: excluded.tsv	• () ×
90: FROGS Filters: abundance.biom	• / X
<u>89: FROGS Filters:</u> sequences.fasta	• / ×

If Filters fields are « Apply » so you have to fill at one field. Otherwise, galaxy become red !

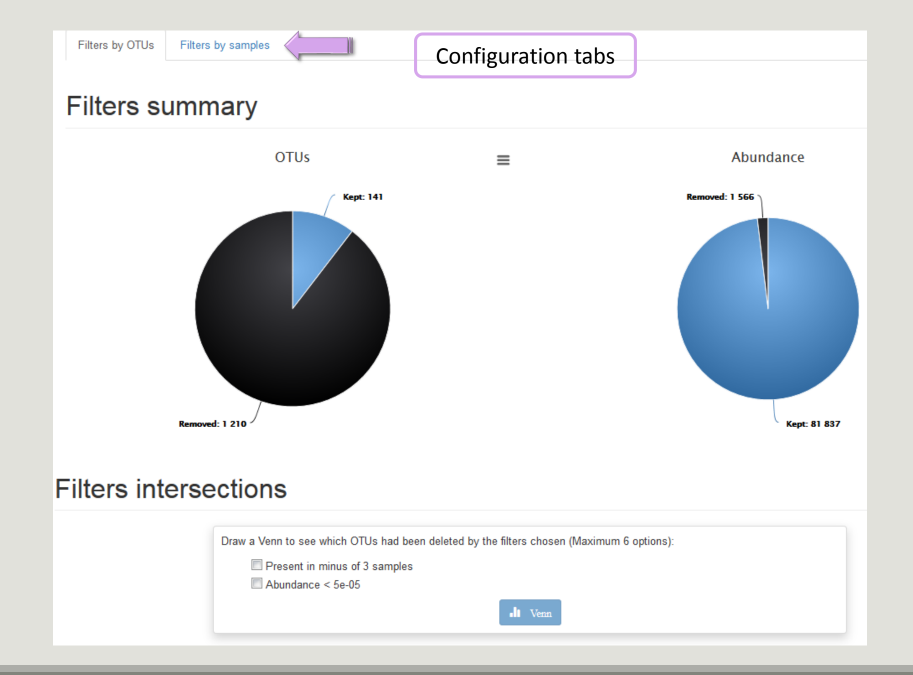
Options

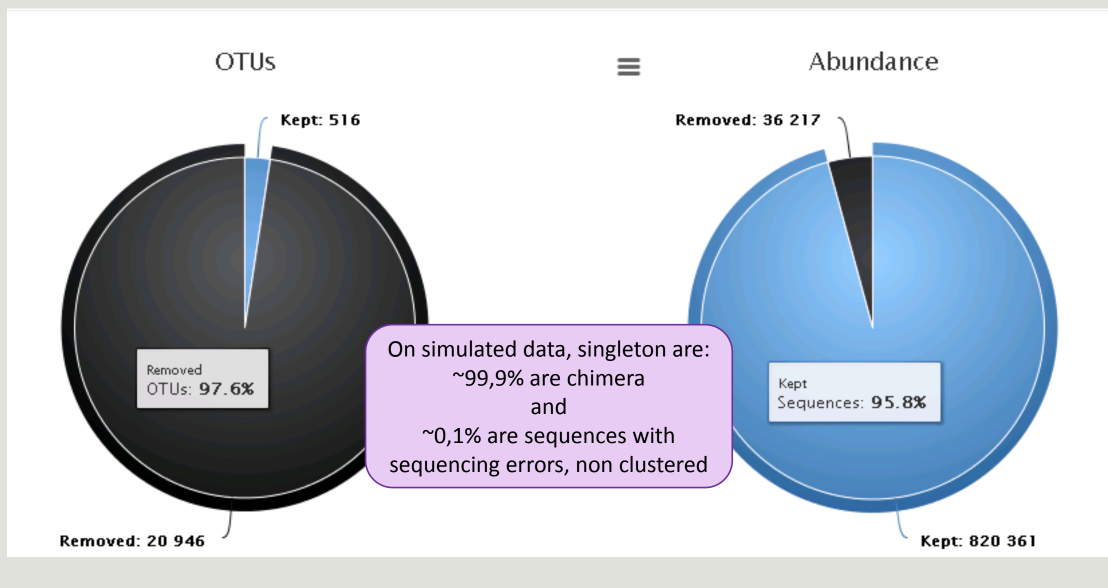
.

#### MiSeq contiged

## Exercise 6

- 1. What are the output files of "Filters"?
- 2. Explore "FROGS Filter : report.html" file.
- 3. How many OTUs have you removed ?
- 4. Build the Venn diagram on the two filters.
- 5. How many OTUs have you removed with each filter "abundance > 0.005%", "Remove OTUs that are not present at least in 3 samples"?
- 6. How many OTUs do they remain ?
- 7. Is there a sample more impacted than the others ?
- 8. To characterize these new OTUs, do not forget to launch "FROGS Cluster Stat" tool, and rename the output HTML file.





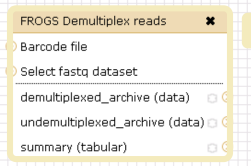
Removing little OTUs (conservation rate =0.005%) and non shared OTU (in less than 2 samples)

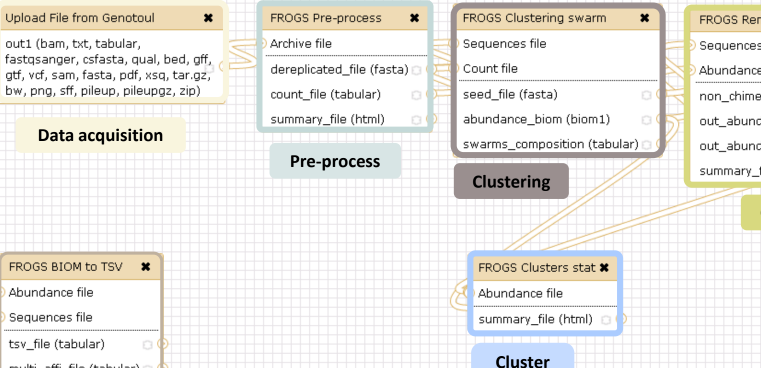
#### Venn on removed OTUs



х

# Affiliation tool





**Statistics** 

Demultiplexing

# FROGS Remove chimera FROGS A Sequences file OTU see Abundance file Abunda non\_chimera\_fasta (fasta) Image: Control of the summar out\_abundance\_biom (biom1) Image: Control of the summar out\_abundance\_count (tabular) Image: Chimera FROGS Filters Sequences file Abundance file Image: Chimera Guide file Image: Chimera Abundance file Image: Chimera Image: Chimera Image: Chimera

×

8(

output\_fasta (fasta)

output\_biom (biom1)

output\_summary (html)

output\_excluded (tabular) 🖸

**Convert to TSV** 

-multi\_affi\_file (tabular) 🖂 🌗

	Affiliation		FROGS Affiliation OTU Step 4 in metagenomics analysis : Taxonomic affiliation of
sur	mmary (html)	8	
bio	m_affiliation (biom	1) 🖂 🤇	
Abr	undance file		
ОТ	U seed sequence		
FR	OGS Affiliation OTU	×	

FROGS Affiliation OTU Step 4 in metagenomics analysis :	Taxonomic affiliation	of each OTU's seed by RDPtools and BLAST	▼ Options
(Galaxy Version 0.8.0)			
Using reference database			
silva123 165 OR	silva128 16S		•
Select reference from the list	silva128 18S		
Also perform RDP assignation?	silva128 23S		
	silva123 16S		
Yes No Optional	silva123 23S	reform it also with PDD classifier (default No)	
Taxonomy affiliation will be perform thanks to Blast. This o	silva123 18S	erform it also with RDP classifier (default No)	
OTU seed sequence	greengenes13_5		
🗋 🔁 🗅 17: FROGS Filters: sequences.fasta	midas_S123_2.1.3		-
OTU sequences (format: fasta).	midas_S119_1.20		
Abundance file	pr2_gb203_4.5		
18: FROGS Filters: abundance.biom			-
OTU abundances (format: BIOM).			
✓ Execute			

## 1 Cluster = 2 affiliations

**Double Affiliation vs** SILVA 123 (for 16S, 18S or 23S), SILVA 119 (for 18S) or Greengenes with :

1. RDPClassifier\* (Ribosomal Database Project): one affiliation with bootstrap, on each taxonomic subdivision.

Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Pseudobutyrivibrio(80); Pseudobutyrivibrio xylanivorans (80)

2. NCBI Blastn+\*\* : all identical Best Hits with identity %, coverage %, e-value, alignment length and a special tag "**Multi-affiliation**".

Bacteria; Firmicutes; Clostridia; Clostridiales; Lachnospiraceae; Pseudobutyrivibrio; Pseudobutyrivibrio ruminis; Pseudobutyrivibrio xylanivorans Identity: 100% and Coverage: 100%

> \* Appl. Environ. Microbiol. August 2007 vol. 73 no. 16 5261-5267. doi : 10.1128/AEM.00062-07 Naïve Bayesian Classifier for Rapid Assignment of rRNA Sequences into the New Bacterial Taxonomy. Qiong Wang, George M.Garrity, James M. Tiedje and James R. Cole

## Affiliation Strategy of FROGS

Blastn+ with "Multi-affiliation" management

V3 – V4	Bacteria   Firmicutes   Clostridia   Clostridiales   Lachnospiraceae   Pseudobutyrivibrio   16S unknown species
V3 – V4	Bacteria   Firmicutes   Clostridia   Clostridiales   Lachnospiraceae   Pseudobutyrivibrio   16S Butyrivibrio fibrisolvens
V3 – V4	Bacteria   Firmicutes   Clostridia   Clostridiales   Lachnospiraceae   Pseudobutyrivibrio   16S rumen bacterium 8   9293-9
V3 – V4	Bacteria   Firmicutes   Clostridia   Clostridiales   Lachnospiraceae   Pseudobutyrivibrio   16S Pseudobutyrivibrio xylanivorans
V3 – V4	Bacteria   Firmicutes   Clostridia   Clostridiales   Lachnospiraceae   Pseudobutyrivibrio   16S Pseudobutyrivibrio ruminis

5 identical blast best hits on SILVA 123 databank

## Affiliation Strategy of FROGS

Blastn+ with "Multi-affiliation" management

V3 – V4	Bacteria   Firmicutes   Clostridia   Clostridiales   Lachnospiraceae   Pseudobutyrivibrio   16S unknown species
V3 – V4	Bacteria   Firmicutes   Clostridia   Clostridiales   Lachnospiraceae   Pseudobutyrivibrio   16S Butyrivibrio fibrisolvens
V3 – V4	Bacteria   Firmicutes   Clostridia   Clostridiales   Lachnospiraceae   Pseudobutyrivibrio   16S rumen bacterium 8   9293-9
V3 – V4	Bacteria   Firmicutes   Clostridia   Clostridiales   Lachnospiraceae   Pseudobutyrivibrio   16S Pseudobutyrivibrio xylanivorans
V3 – V4	Bacteria   Firmicutes   Clostridia   Clostridiales   Lachnospiraceae   Pseudobutyrivibrio   16S Pseudobutyrivibrio ruminis

**FROGS Affiliation:** Bacteria | Firmicutes | Clostridia | Clostridiales | Lachnospiraceae | Pseudobutyrivibrio | **Multi-affiliation** 

# Your Turn! – 7

LAUNCH THE « FROGS AFFILIATION » TOOL



## Exercise 7.1

Go to « MiSeq contiged » history

Launch the « FROGS Affiliation » tool with

- SILVA 123 or 128 16S database
- FROGS Filters abundance biom and fasta files (after swarm d1d3, remove chimera and filter low abundances)
- $\rightarrow$  objectives :
  - understand abundance tables columns
  - understand the BLAST affiliation

#### FROGS Affiliation OTU X

OTU seed sequence

Abundance file

biom\_affiliation (biom1) 🖂 🤇

summary (html)

#### Affiliation

Using reference data	base	
silva123 16S		
Select reference from t	the list	
Also perform RDP ass	signation?	
Yes No Taxonomy affiliation wi	ill be perform thanks to Blast. This option allow you to perform it also with RDP classifier (default No)	
OTU seed sequence		
C 2 C 17: F	FROGS Filters: sequences.fasta	
OTU sequences (forma	at: fasta).	
Abundance file		
18: F	ROGS Filters: abundance.biom	
	nat: BIOM).	



## Exercise 7.1

- 1. What are the « FROGS Affiliation » output files ?
- 2. How many sequences are affiliated by BLAST ?
- 3. Click on the « eye » button on the BIOM output file, what do you understand ?
- Use the Biom\_to\_TSV tool on this last file and click again on the "eye" on the new output generated.
   What do the columns ?

What do the columns

What is the difference if we click on case or not ? What consequence about weight of your

file ?

FROGS BIOM to TSV Converts a BIOM file in TSV file. (Galaxy Version 2.1.0)
Abundance file
C 22: FROGS Affiliation OTU: affiliation.biom
The BIOM file to convert (format: BIOM).
Sequences file
C 2 Nothing selected
The sequences file (format: fasta). If you use this option the sequences will be add in TSV.
Extract multi-alignments
Yes No
If you have used FROGS affiliation on your data, you can extract information about multiple alignements in a second TSV.
✓ Execute

#### Tools

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FROGS - FIND RAPIDLY OTU WITH GALAXY SOLUTION

#### FROGS pipeline

FROGS Upload archive from your computer

FROGS Demultiplex reads Split by samples the reads in function of inner barcode.

FROGS Pre-process Step 1 in metagenomics analysis: denoising and dereplication.

FROGS Clustering swarm Step 2 in metagenomics analysis : clustering.

FROGS Remove chimera Step 3 in metagenomics analysis : Remove PCR chimera in each sample.

<u>FROGS Filters</u> Filters OTUs on several criteria.

FROGS Affiliation OTU Step 4 in metagenomics analysis : Taxonomic affiliation of each OTU's seed by RDPtools and BLAST

FROGS BIOM to TSV Converts a BIOM file in TSV file.

FROGS Clusters stat Process some metrics on clusters.

FROGS Affiliations stat Process some metrics on taxonomies.

<u>FROGS BIOM to std BIOM</u> Converts a FROGS BIOM in fully compatible BIOM.

FROGS Abundance normalisation



## Exercise 7.1

5. Understand Blast affiliations - Cluster\_2388 (affiliation from silva 123)

blast_subject	blast_evalue	blast_len	blast_perc_q uery_covera ge	blast_perc_id entity	blast_taxonomy
JN880417.1.1422	0.0	360	88.88	99.44	Bacteria;Planctomycetes;Planctomycetacia;Pl anctomycetales;Planctomycetaceae;Telmatoc ola;Telmatocola sphagniphila

## Blast JN880417.1.1422 vs our OTU

#### OTU length : 405

#### Excellent blast but no matches at the beginning of OTU.

Telmatocola sphagniphila strain SP2 16S ribosomal RNA gene, partial sequence Sequence ID: ref[NR 118328.1 Length: 1422 Number of Matches: 1

Range 1: 375 to 734 GenBank Graphics Vext Match 🛦 Prev					
Score		Expect	Identities	Gaps	Strand
654 b	its(35	4) 0.0	358/360(99%)	0/360(0%)	Plus/Plus
Query	46	CGCGTGCGCGATGAAG	GCCTTCGGGTTGTAAAGCG		GAAACCT 105
Sbjct	375		SCCTTCGGGTTGTAAAGCG		GAAACTT 434
Query	106		GCTCGGGCTAAGTTTGTGC		
Sbjct	435		GCTCGGGCTAAGTTTGTGC		
Query	166		ATCACTGGGCATAAAGGGC		
Sbjct	495		ATCACTGGGCATAAAGGGC		
Query	226	GTGAAATACTTCAGCT	CAACTGGAGAACTGCCTCG	GATACTGGGAATCTCGAG	TAATGTA 285
Sbjct	555	GTGAAATACTTCAGCT	CAACTGGAGAACTGCCTCG	GATACTGGGAATCTCGAG	TAATGTA 614
Query	286		TGGTGGAGCGGTGAAATG		
Sbjct	615	GGGGCACGTGGAACGG	TGGTGGAGCGGTGAAATG	CGTTGATATCAGTCGGA	ACTCCGGT 674
Query	346	GGCGAAGGCGATGTGC	GGACATTTACTGACGCTG	AGGCGCGAAAGCCAGGGG	AGCAAAC 405
Sbjct	675	GGCGAAGGCGATGTGC	rggacatttactgacgctg	AGGCGCGAAAGCCAGGG	AGCAAAC 734

#### Telmatocola sphagniphila strain SP2 16S ribosomal RNA gene, partial sequence

NCBI Reference Sequence: NR\_118328.1

FASTA Graphics

#### <u>Go to:</u> 🖂

LOCUS	NR_118328 1422 bp rRNA linear BCT 03-FEB-2015
DEFINITION	Telmatocola sphagniphila strain SP2 16S ribosomal RNA gene, partial
ACCESSIO VERSION DBLINK	NR_118328 IIII:645321338 Project: 33175
DDDIRK	BioProject: PRJNA33175
KEYWORDS	RefSeq.
SOURCE	Telmatocola sphagniphila
ORGANISM	Telmatocola sphagniphila
	Bacteria; Planctomycetes; Planctomycetia; Planctomycetales;
	Planctomycetaceae.
REFERENCE	1 (bases 1 to 1422)
AUTHORS	Kulichevskaya,I.S., Serkebaeva,Y.M., Kim,Y., Rijpstra,W.I.,
	Damste,J.S., Liesack,W. and Dedysh,S.N.
TITLE	Telmatocola sphagniphila gen. nov., sp. nov., a novel dendriform
	planctomycete from northern wetlands
JOURNAL	Front Microbiol 3, 146 (2012)
PUBMED	22529844
REMARK	Publication Status: Online-Only
	2 (bases 1 to 1422)
CONSRTM	
TITLE	Direct Submission
JOURNAL	Submitted (28-APR-2014) National Center for Biotechnology
	Information, NIH, Bethesda, MD 20894, USA
REFERENCE	3 (bases 1 to 1422)
AUTHORS	Dedysh, S.N.
TITLE	Direct Submission
JOURNAL	Submitted (20-OCT-2011) Winogradsky Institute of Microbiology RAS,
CONVENT	Prospect 60-Letya Octyabrya 7/2, Moscow 117312, Russia
COMMENT	REVIEWED <u>REFSEQ</u> : This record has been ourgoed by mail staff. The
	reference sequence is identical to JN880417:1-1422.

## Blast columns

### OTU\_2 seed has a best BLAST hit with the reference sequence AJ496032.1.1410

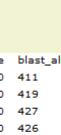
## The reference sequence taxonomic affiliation is this one.

#blast_taxonomy	blast_subject	blast_perc_identity	blast_perc_query_coverage	blast_evalue	blast_aln_length
Bacteria; Actinobacteria; Actinobacteria; Bifidobacteriales; Bifidobacteriaceae; Metascardovia; Multi-affiliation	multi-subject	100.0	100.0	0.0	411
Bacteria;Fibrobacteres;Fibrobacteria;Fibrobacterales;Fibrobacteraceae;Fibrobacter;Fibrobacter succinogenes	AJ496032.1.1410	100.0	100.0	0.0	419
Bacteria;Firmicutes;Bacilli;Bacillales;Staphylococcaceae;Nosocomiicoccus;Nosocomiicoccus ampullae	EU240886.1.1502	100.0	100.0	0.0	427
Bacteria; Proteobacteria; Gammaproteobacteria; Pseudomonadales; Moraxellaceae; Psychrobacter; Psychrobacter immobilis	U39399.1.1477	100.0	100.0	0.0	426
Bacteria;Thermotogae;Thermotogae;Thermotogales;Thermotogaceae;Petrotoga;Petrotoga miotherma	FR733705.1.1499	100.0	100.0	0.0	419
${\tt Bacteria}; {\tt Proteobacteria}; {\tt Alphaproteobacteria}; {\tt Rhizobiales}; {\tt Phyllobacteriaceae}; {\tt Pseudahrensia}; {\tt Pse$	GU575117.1.1441	100.0	100.0	0.0	401
Bacteria;Bacteroidetes;Cytophagia;Cytophagales;Cytophagaceae;Persicitalea;Persicitalea jodogahamensis	multi-subject	100.0	100.0	0.0	421
${\tt Bacteria}; {\tt Proteobacteria}; {\tt Deltaproteobacteria}; {\tt Bdellovibrionales}; {\tt Bdellovibrionaceae}; {\tt Bdellovibrio}; {\tt Multi-affiliation}; {\tt Multi-af$	multi-subject	100.0	100.0	0.0	404

#### **Convert to TSV**

FROGS BIOM to TSV
Abundance file
Sequences file
tsv_file (tabular) 🛛 🔅 🤇
multi_affi_file (tabular) 🖂 🤇

#### Evaluation variables of BLAST



Does

Kennard Play

Classical

Guitar

Songs?

Or Folk

DOMAIN

Kingdom Phylum

Class

Order

Family

Genus Species

## Focus on "Multi-"

#### (affiliation from silva 123)

#### Observe line of Cluster 1 inside abundance.tsv and multi\_hit.tsv files, what do you conclude ?

#blast_taxonomy	blast_subject	blast_perc_identity	blast_perc_query_coverage	blast_evalue	blast_aln_length
Bacteria; Actinobacteria; Actinobacteria; Bifidobacteriales; Bifidobacteriaceae; Metascardovia; Multi-affiliation	multi-subject	100.0	100.0	0.0	411
Bacteria;Fibrobacteres;Fibrobacteria;Fibrobacterales;Fibrobacteraceae;Fibrobacter;Fibrobacter succinogenes	AJ496032.1.1410	100.0	100.0	0.0	419
Bacteria;Firmicutes;Bacilli;Bacillales;Staphylococcaceae;Nosocomiicoccus;Nosocomiicoccus ampullae	EU240886.1.1502	100.0	100.0	0.0	427
Bacteria; Proteobacteria; Gammaproteobacteria; Pseudomonadales; Moraxellaceae; Psychrobacter; Psychrobacter immobilis	U39399.1.1477	100.0	100.0	0.0	426
Bacteria; Thermotogae; Thermotogae; Thermotogales; Thermotogaceae; Petrotoga; Petrotoga miotherma	FR733705.1.1499	100.0	100.0	0.0	419
Bacteria ; Proteobacteria ; Alpha proteobacteria ; Rhizobiales ; Phyllobacteria ceae ; Pseudahrensia ; Pseudahrensia aquimaris a substantia a subs	GU575117.1.1441	100.0	100.0	0.0	401
Bacteria;Bacteroidetes;Cytophagia;Cytophagales;Cytophagaceae;Persicitalea;Persicitalea jodogahamensis	multi-subject	100.0	100.0	0.0	421
Bacteria ; Proteobacteria ; Delta proteobacteria ; Bdellovibrionales ; Bdellovibrionaceae ; Bdellovibrio ; Multi-affiliation = (Content of the second seco	multi-subject	100.0	100.0	0.0	404

Cluster\_1 has 5 identical blast hits, with different taxonomies as the species level

## Focus on "Multi-"

(affiliation from silva 123)

Observe line of Cluster 11 inside abundance.tsv and multi\_hit.tsv files, what do you conclude ?

Bacteria; Proteobacteria; Alphaproteobacteria; Caulobacterales; Hyphomonadaceae; Henriciella; Henriciella marina	multi-subject	100.0 100	0.0
------------------------------------------------------------------------------------------------------------------	---------------	-----------	-----

Cluster\_11 has 2 identical blast hits, with identical species but with different strains (strains are not written in our data)

## Focus on "Multi-"

(affiliation from silva 123)

Observe line of Cluster 43 inside abundance.tsv and multi_hit.tsv files, what do you conclude ?							
Bacteria;Firmicutes;	;Negativicutes;Selenomonadales;Veillonellaceae;Multi-affiliation;Multi-affiliation	multi-subject	99.3	100.0			
Cluster_43	Bacteria;Firmicutes;Negativicutes;Selenomonadales;Veillonellaceae;Selenomonas 3;unkn	own species	JQ	447821.1.1420			
Cluster_43	Bacteria;Firmicutes;Negativicutes;Selenomonadales;Veillonellaceae;Centipeda;Centipeda	a periodontii	AJ	010963.1.1494			



Cluster\_43 has 2 identical blast hits, with different taxonomies at the genus level

## Back on Blast parameters

#blast_taxonomy	blast_subject	blast_perc_identity	blast_perc_query_coverage	blast_evalue	blast_aln_length
Bacteria; Actinobacteria; Actinobacteria; Bifidobacteriales; Bifidobacteriaceae; Metascardovia; Multi-affiliation	multi-subject	100.0	100.0	0.0	411
Bacteria;Fibrobacteres;Fibrobacteria;Fibrobacterales;Fibrobacteraceae;Fibrobacter;Fibrobacter succinogenes	AJ496032.1.1410	100.0	100.0	0.0	419
Bacteria;Firmicutes;Bacilli;Bacillales;Staphylococcaceae;Nosocomiicoccus;Nosocomiicoccus ampullae	EU240886.1.1502	100.0	100.0	0.0	427
Bacteria ; Proteobacteria ; Gamma proteobacteria ; Pseudomonadales ; Moraxellaceae ; Psychrobacter ; Psychrobacter immobilis and the second	U39399.1.1477	100.0	100.0	0.0	426
Bacteria;Thermotogae;Thermotogae;Thermotogales;Thermotogaceae;Petrotoga;Petrotoga miotherma	FR733705.1.1499	100.0	100.0	0.0	419
Bacteria ; Proteobacteria ; Alpha proteobacteria ; Rhizobiales ; Phyllobacteriaceae ; Pseudahrensia ; Pseudahrensia a quimaris a substitution of the second statement of the	GU575117.1.1441	100.0	100.0	0.0	401
Bacteria;Bacteroidetes;Cytophagia;Cytophagales;Cytophagaceae;Persicitalea;Persicitalea jodogahamensis	multi-subject	100.0	100.0	0.0	421
${\tt Bacteria} \\ {\tt Proteobacteria} \\ {\tt Delta proteobacteria} \\ {\tt Bdellovibrionales} \\ {\tt Bdellovibrionaceae} \\ {\tt Bdellovibrio} \\ {\tt Multi-affiliation} \\ {\tt Multi-affiliat$	multi-subject	100.0	100.0	0.0	404

Evaluation variables of BLAST

## Blast variables : e-value

The Expect value (E) is a parameter that describes the number of hits one can "expect" to see by chance when searching a database of a particular size.

The lower the E-value, or the closer it is to zero, the more "significant" the match is.

## Blast variables : blast\_perc\_identity

Identity percentage between the Query (OTU) and the subject in the alignment (length subject = 1455 bases)

Score		Expect	Identities	Gaps	Strand		
760 bit	s(411	) 0.0	411/411(100%)	0/411(0%)	Plus/P	lus	
~	-	111111111111111111	ATGGGGGGAACCCTGATG 			60 390	
Query Sbjct		1111111111111111111	CGCTTTTAATTGGGAGCAA			120 450	Query length = 411 Alignment length = 4
~ 1		111111111111111111	TAACTACGTGCCAGCAGCCC		1111111	180 510	0 mismatch
~ -		1111111111111111111	CGTAAAGAGCTCGTAGGCC			240 570	-> 100% identity
~ -			GATTTGCGCTGGGTACGGG 			300 630	
~ -		111111111111111111	ACGGTGGAATGTGTAGATA                      ACGGTGGAATGTGTAGATA			360 690	
~		1111111111111111111	GACTGACGCTGAGGAGCGAA 		411 741		

411

## Blast variables : blast\_perc\_identity

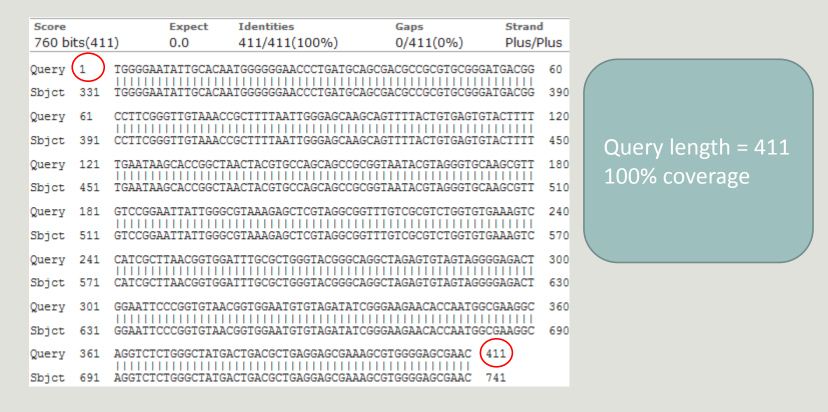
Identity percentage between the Query (OTU) and the subject in the alignment (length subject = 1455 bases)

Score		Expect	Identities	Gaps	Strand	
614 bi	ts(332)	5e-172	385/411(94%)	5/411(1%)	Plus/Plus	
Query	1	TGGGGAATATTGCAC	AATGGGGGGGAACCCTGATGCA	GCGACGCCGCGTGCGG		60
Sbjct	140728		AATGGGCGAAAGCCTGATGCA			140787
Query	61		CCGCTTTTAATTGGGAGCAAG		GTACTTTT	120
Sbjct	140788		CCGCTTTTGATTGGGAGCAAG			140842
Query	121		IAACTACGTGCCAGCAGCCGC			180
Sbjct	140843		TAACTACGTGCCAGCAGCCGC			140902
Query	181		GCGTAAAGAGCTCGTAGGCGG			240
Sbjct	140903		GCGTAAAGRGCTCGTAGGCGG			140962
Query	241		GATTTGCGCTGGGTACGGGCA		GGGAGACT	300
Sbjct	140963		GATCTGCGCCGGGTACGGGCG			141022
Query	301		ACGGTGGAATGTGTAGATATC			360
Sbjct	141023		ACGGTGGAATGTGTAGATATC			141082
Query	361	AGGTCTCTGGGCTAT	GACTGACGCTGAGGAGCGAAA		411	
Sbjct	141083		IACTGACGCTGAGGAGCGAAA		141133	

Query length = 411 Alignment length = 411 26 mismatches (gaps included) -> 94% identity

## Blast variables : blast\_perc\_query\_coverage

#### Coverage percentage of alignment on query (OTU)



## Blast variables : blast-length

Length of alignment between the OTUs = "Query" and "subject" sequence of database

	Coverage %	Identity %	Length alignment
OTU1	100	98	400
OTU2	100	98	500

FROGS Affiliation OTU Step 4 in metagenomics analy	rsis : Taxonomic affiliation of each OTU's seed by RDPtools and BLAST
(Galaxy Version 0.8.0)	
Using reference database	
silva123 16S	
Select reference from the list	
Also perform RDP assignation? Yes No Taxonomy amiliation will be perform thanks to Blast. If OTU seed sequence	Optional and <u>not</u> in our guideline
17: FROGS Filters: sequences.fasta	
OTU sequences (format: fasta).  Abundance file  18: FROGS Filters: abundance.biom	Who have already used
OTU abundances (format: BIOM).	RDP previously ?

FROGS Affiliation OTU

biom\_affiliation (biom1) 🗇

Affiliation

0

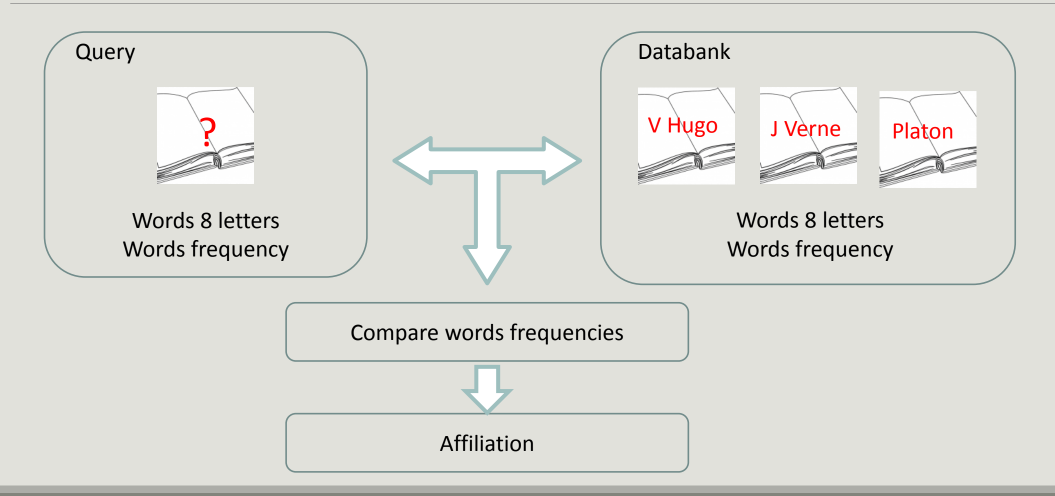
OTU seed sequence

Abundance file

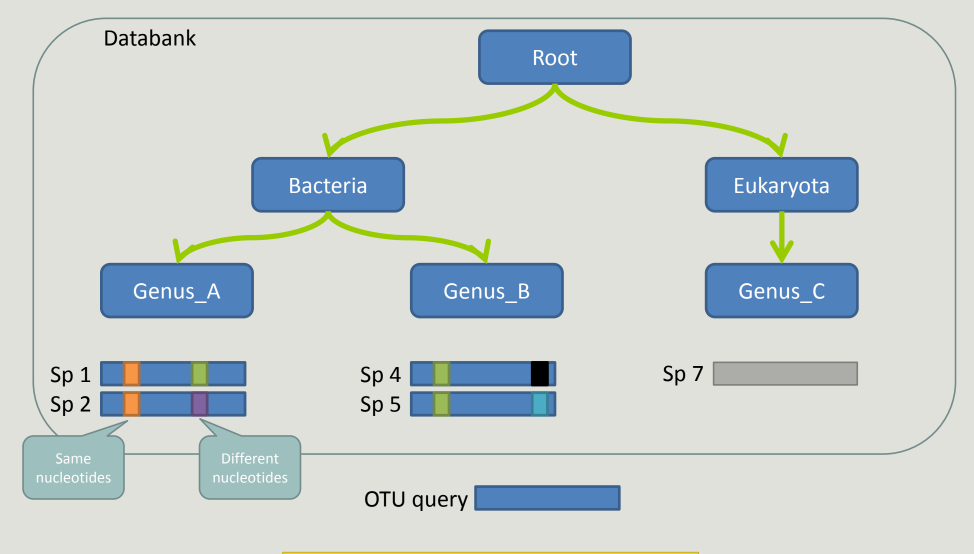
summary (html)

Escape RDP explanation

## How works RDP ?

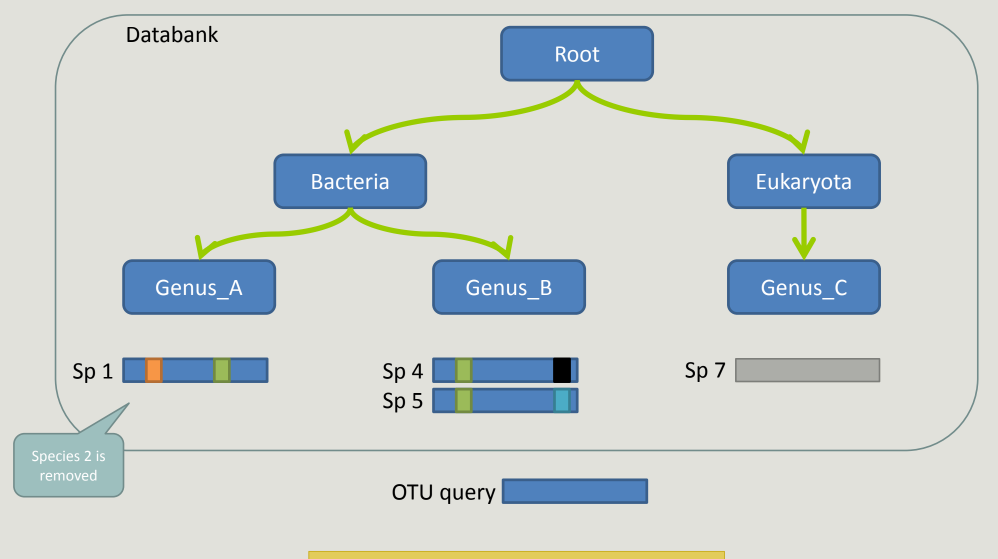


## How works RDP ?

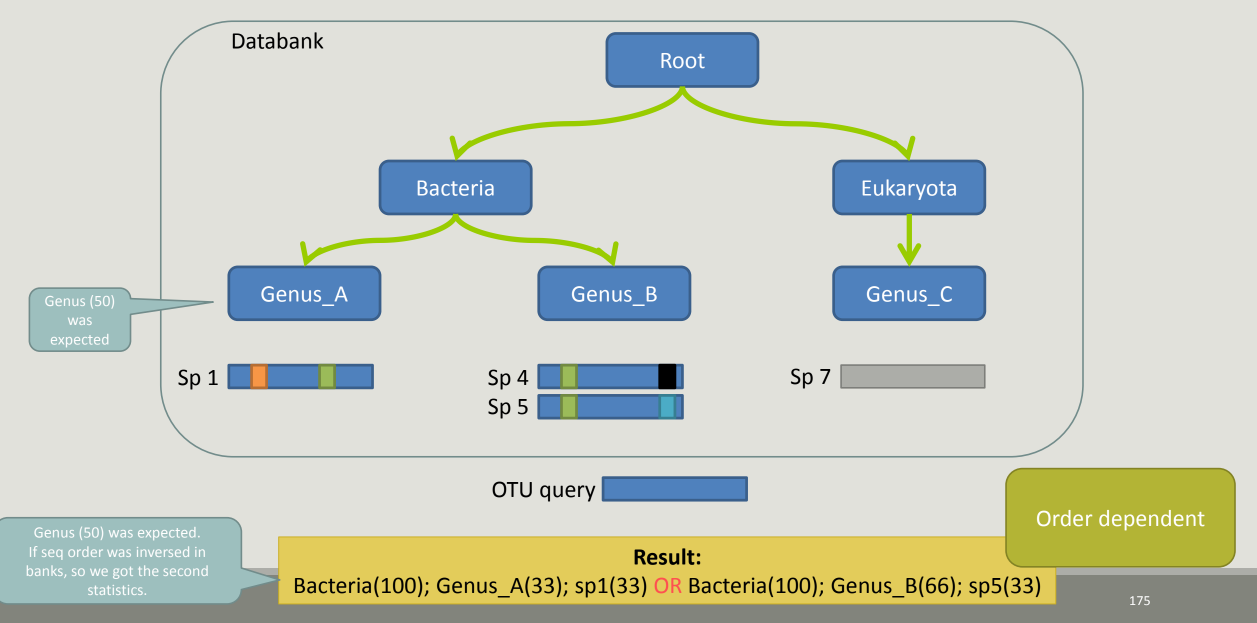


Result: Bacteria(100) ; Genus\_A(50) ; Sp1(25)

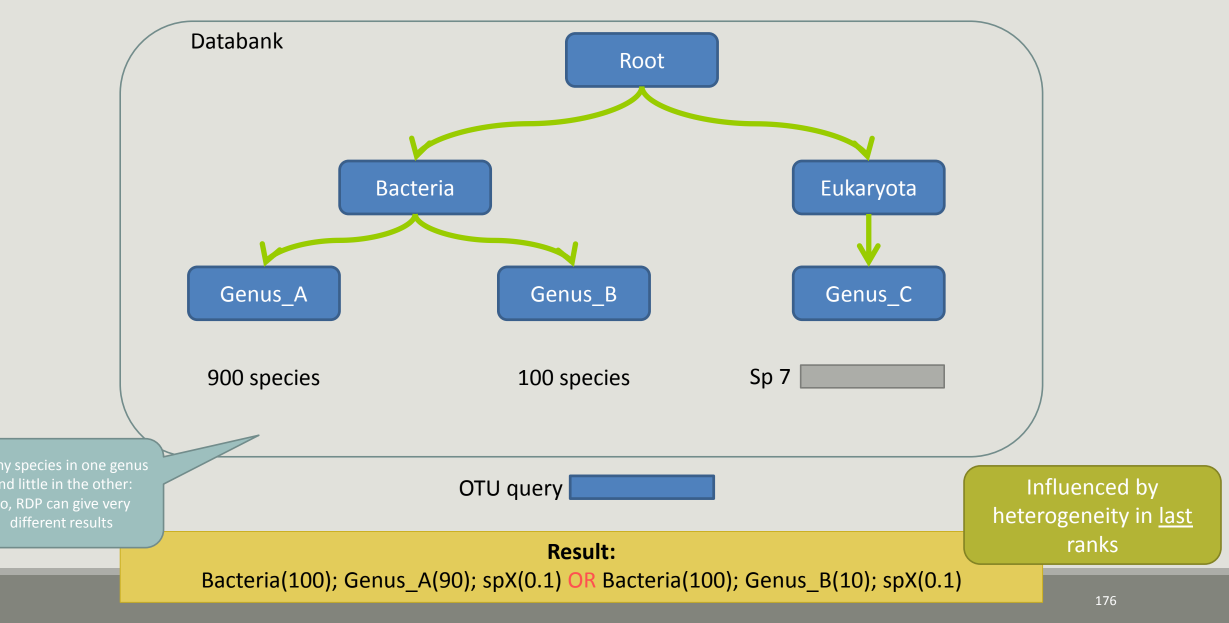
## The dysfunctions of RDP ?



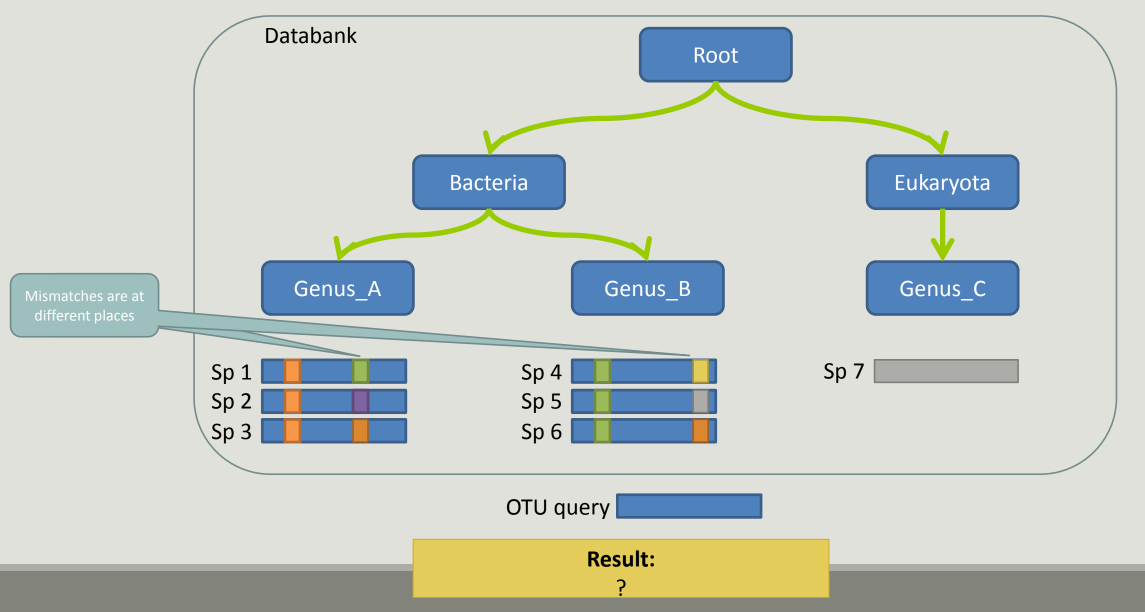
## The dysfunctions of RDP n°1?



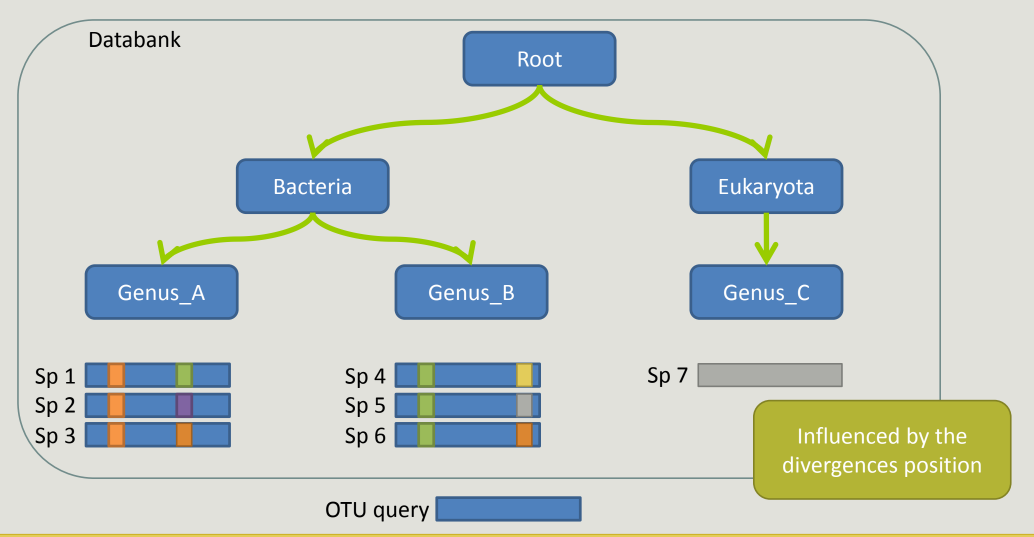
## The dysfunctions of RDP n°2 ?



## The dysfunctions of RDP n°3 ?

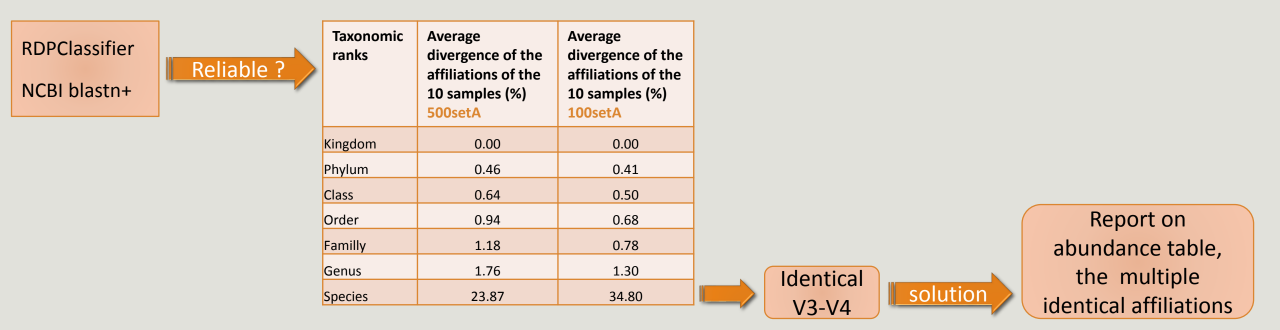


## The dysfunctions of RDP n°3 ?



Si le mismatch se fait sur un mot très "significatif" dans le profil de k-mers, RDP ne tombera que rarement sur l'espèce lors du bootstrap. Avec une même distance d'édition (2 mismatchs) on peut donc avoir une grande différence de bootstrap pour peu que le mot affecté soit important dans le profil. 178

# Divergence on the composition of microbial communities at the different taxonomic ranks



	Only one best	hit			Multiple best	hit
Taxonomic ranks	Average divergence of the affiliations of the 10 samples (%) 500setA	Average divergence of the affiliations of the 10 samples (%) 100setA		Taxonomic ranks	Median divergence of the affiliations of the 10 samples (%) 500setA	Median divergence of the affiliations of the 10 samples (%) 100setA
Kingdom	0.00	0.00		Kingdom	0.00	0.00
Phylum	0.46	0.41		Phylum	0.46	0.41
Class	0.64	0.50		Class	0.64	0.50
Order	0.94	0.68		Order	0.93	0.68
Familly	1.18	0.78		Familly	1.17	0.78
Genus	1.76	1.30		Genus	1.60	1.00
Species	23.87	34.80	Species		6.63	5.75
		W FROGS	ith the S guide		Median divergence of the affiliations of the 10 samples (%) 500setA filter: 0.005% - 505 OTUs	Median divergence of the affiliations of the 10 samples (%) 100setA filter: 0.005% - 100 OTUs
				Kingdom	0.00	0.00
				Phylum	0.38	0.38
				Class	0.57	0.48
				Order	0.81	0.64
				Familly	1.08	0.74
				Genus	1.43	0.76

Species

1.53

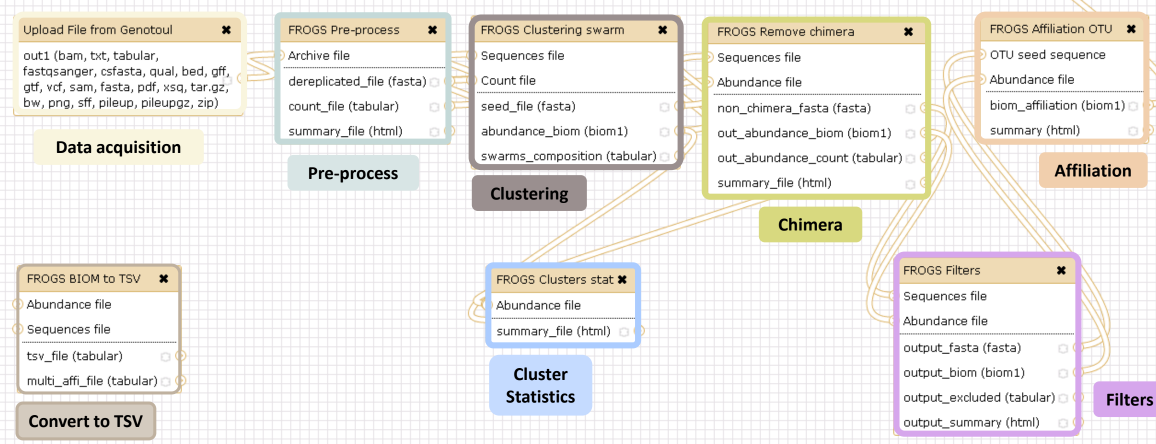
0.78

# Careful: Multi hit blast table is non exhaustive !

- Chimera (multiple affiliation)
- V3V4 included in others
- Missed primers on some 16S during database building

# Affiliation Stat





FROGS Affiliations stat Process some metrics on taxonomies. (Galaxy Version 1.1.0)	✓ Options	FROGS Affiliations stat Process some metrics on taxonomies. (Galaxy Version 1.1.0)	✓ Options
FROGS Affiliations stat Process some metrics on taxonomies. (Galaxy Version 1.1.0)         Abundance file	Options	FROGS Affiliations stat Process some metrics on taxonomies. (Galaxy Version 1.1.0)         Abundance file         22: FROGS Affiliation OTU: affiliation.biom         OTUs abundances and affiliations (format: BIOM).         Rarefaction ranks         Class Order Family Genus Species         The ranks that will be evaluated in rarefaction. Each rank is separated by one space.         Affiliation processed         FROGS rdp         Subset the type of affiliation processed. If your affiliation has been processed with an external tool: use 'Custom'.	Options
Taxonomy distribution       Alignment distribution		Control   Control Con	
		The metadata tag used in BIOM file to store the alignment OTUs coverage.	

✓ Execute

# Exercise 7.2

## FROGS Affiliations stat (version 1.1.0)

### Abundance file:

17: FROGS Affiliation OTU: affiliation.biom

OTUs abundances and affiliations (format: BIOM).

## Rarefaction ranks:

## **Class Order Family Genus Species**

The ranks that will be evaluated in rarefaction. Each rank is separated by one space.

#### Affiliation processed:

FROGS blast ᅌ

Select the type of affiliation processed. If your affiliation has been processed with an external tool: use 'Custom'.

### Execute

### FROGS Affiliations stat (version 1.1.0)

## Abundance file:

17: FROGS Affiliation OTU: affiliation.biom

OTUs abundances and affiliations (format: BIOM).

## **Rarefaction ranks:**

## **Class Order Family Genus Species**

The ranks that will be evaluated in rarefaction. Each rank is separated by one space.

## Affiliation processed:

# Is it adequate on our data ? Why ?

0

Select the type of affiliation processed. If your affiliation has been processed with an external tool: use 'Custom'.

## Execute

FROGS rdp

 Omega 23: FROGS
 ● Ø ⋈

 Affiliations stat: summary.html

# Exercise 7.2

 $\rightarrow$  objectives :

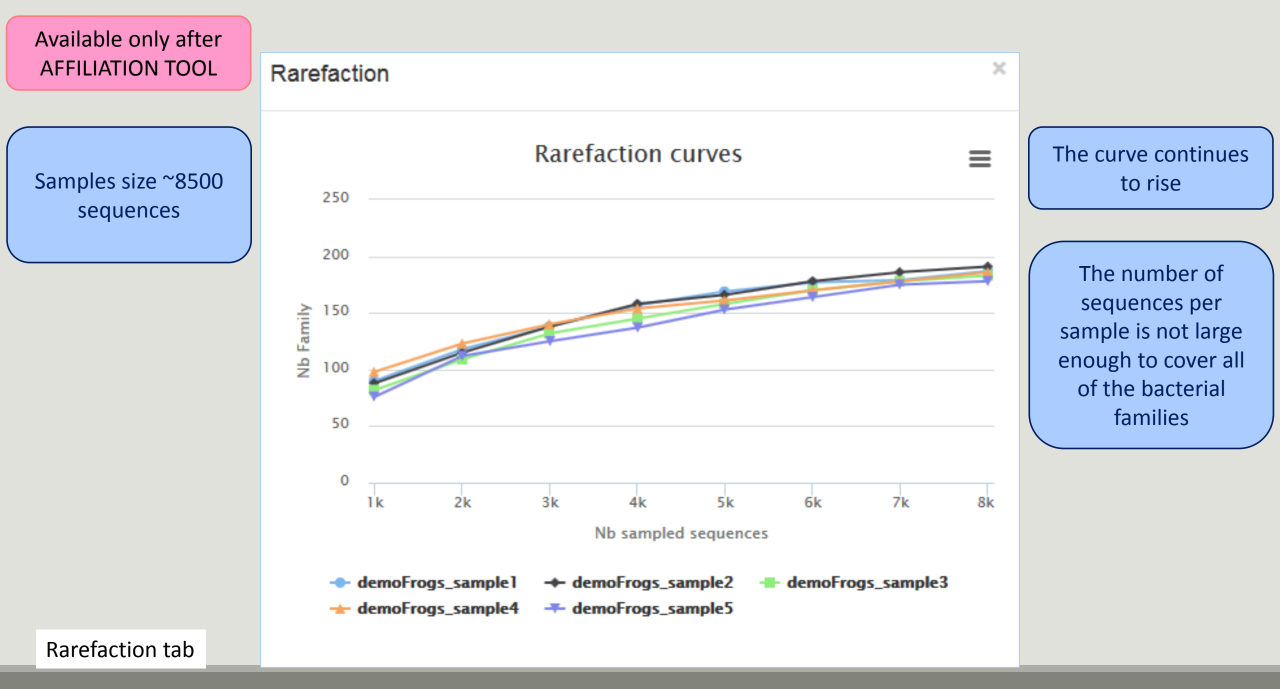
understand rarefaction curve and sunburst

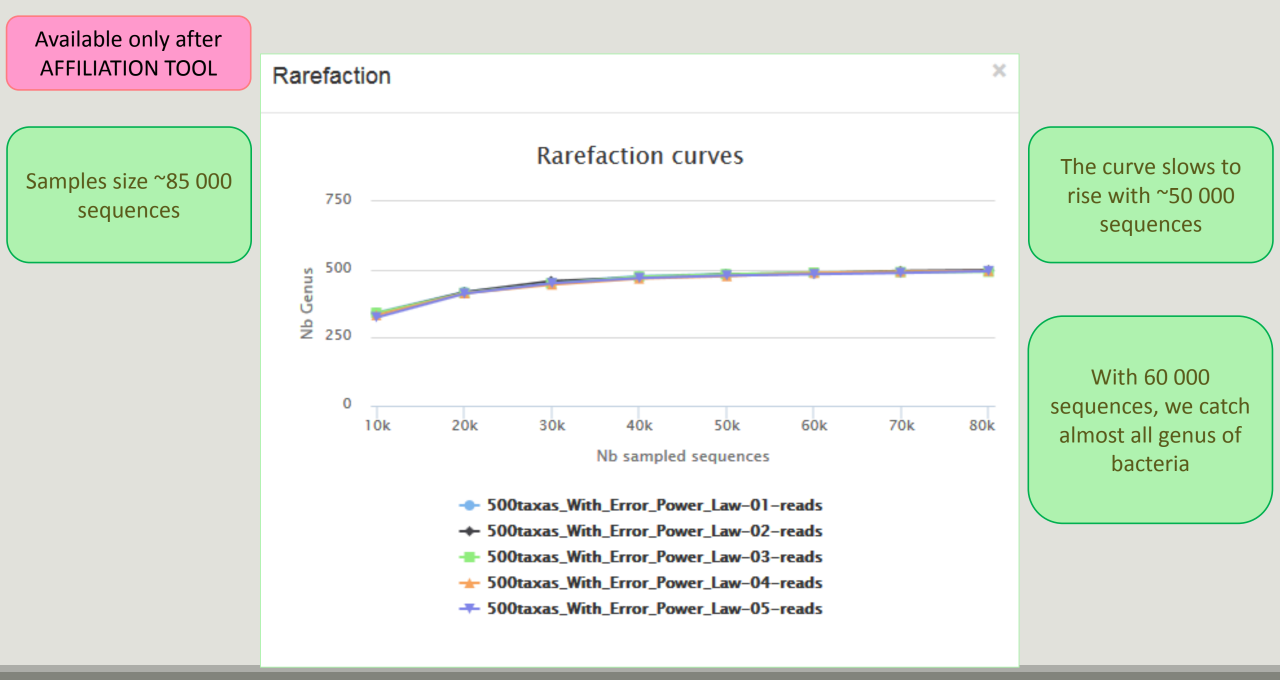
1. Explore the Affiliation stat results on FROGS blast affiliation.

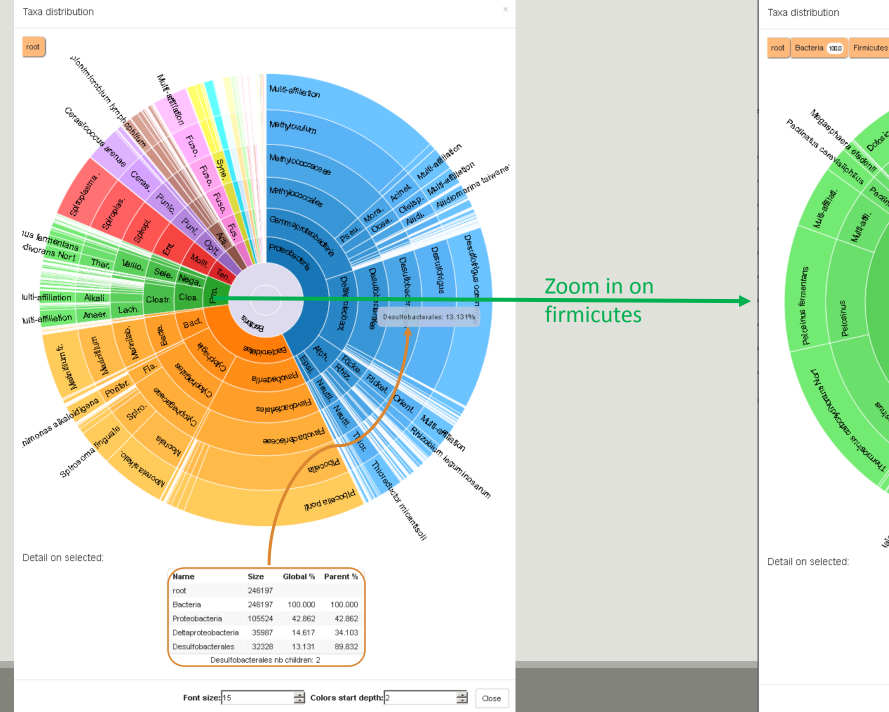
2. What kind of graphs can you generate? What do they mean?

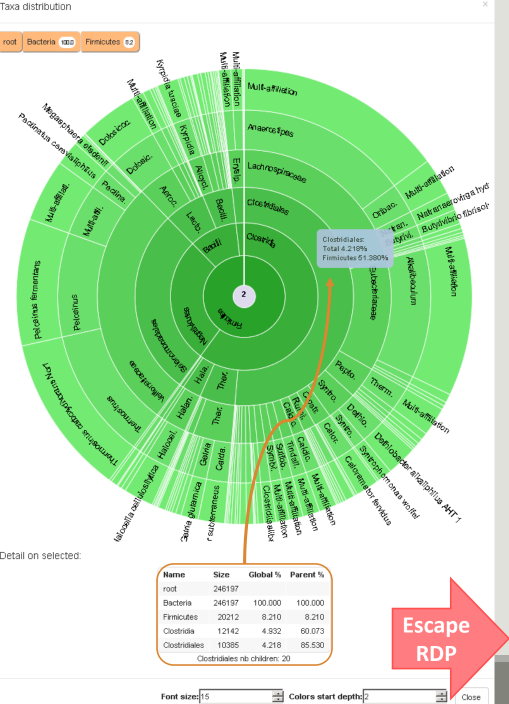
💳 Sigenae - Welcome	e mbernard	Analyze Data Workflow Shared	Data - Visualization -	Admin Help <del>-</del>	User▼				Using 6%
Tools RADSEQ - STACKS RADseqSTACKS	Taxonomy distribution Alignment distribution							History imported: 500WEPL_setA 451.3 MB	2 * 0 •
METHYLATION - BISULFITE Bisulfite BISMARK		alt Display	global distribution					106: FROGS Clusters stat: summary.html	• / X
DEEPTOOLS deepTools							kcsv	<u>105: report_download</u>	• / ×
FROGS - FIND RAPIDLY OTU WITH GALAXY SOLUTION	Show 10 • entries					Search:		103: Vsearch Clusters stat	• / ×
FROGS pipeline	Taxonomies by sample							102: FROGS Affiliations sta summary.html	<u>nt:</u> @
<u>FROGS Upload archive</u> from your computer	Samples	▲ Nb domain Nb phylum	Nb class 🔶 Nb ord	er 🕴 Nb family 🤅	Nb genus	Nb species	Nb sequences 🔶	299.1 KB format: html, database: <u>?</u>	
<u>FROGS Demultiplex reads</u> Split by samples the reads in	☑ 500taxas_With_Error_Power_Law-01-reads	1 29	59 129	243	491	492	81,572	## Application Software: affiliations_stat.py (version:	
function of inner barcode. FROGS Pre-process Step 1 in	00taxas_With_Error_Power_Law-02-reads	1 29	59 130	243	491	492	82,466	Command: /usr/local/bioinfo /src/galaxy-dev/galaxy-dist/t /FROGS/tools/affiliations sta	ools 👘
metagenomics analysis: denoising and dereplication.	500taxas_With_Error_Power_Law-03-reads	1 29	59 130	243	491	493	82,159	input-biom /galaxydata/dat /files/054/dataset_54829.da	tabase
FROGS Clustering swarm Step 2 in metagenomics	500taxas_With_Error_Power_Law-04-reads	1 29	59 130	243	491	492	81,985	output-file /work/galaxy-de 🔚 🛈 🥹	ev/data 🧷 📄
analysis : clustering.	500taxas_With_Error_Power_Law-05-reads	1 29	59 130	241	487	488	82,039	HTML file	
FROGS Remove chimera Step 3 in metagenomics analysis : Remove PCR chimera in each	□ 500taxas_With_Error_Power_Law-06-reads	1 29	59 130	244	493	494	81,758	<u>101: swarm cluster stat</u>	• / ×
sample.	50 (taxas_With_Error_Power_Law-07-reads	1 29	59 130	244	491	492	81,714	100: FROGS BIOM to std	• / ×
<u>FROGS Filters</u> Filters OTUs on several criteria.	500taxas_With_Error_Power_Law-08-reads	1 29	58 129	243	493	494	82,255	BIOM: blast_metadata.tsv	
FROGS Affiliation OTU Step 4 in metagenomics analysis :	500taxas, With_Error_Power_Law-09-reads	1 29	59 130	244	493	494	82,113	<u>99: FROGS BIOM to std</u> <u>BIOM: abundance.biom</u>	• / ×
Taxonomic affiliation of each OTU's seed by RDPtools and BLAST	500taxas_Witb_Error_Power_Law-10-reads	i 29	58 128	240	487	489	82,300	98: FROGS BIOM to TSV: multi_hits.tsv	• / ×
<u>FROGS BIOM to TSV</u> Converts a BIOM file in TSV file.	With selection: Class Display rarefac	tion Display distribution						97: FROGS BIOM to TSV: abundance.tsv	• / ×
FROGS Clusters stat Process some metrics on clusters. FROGS Affiliations stat Process some metrics on taxonomies.	Showing 1 to 10 of 10 entries					Pre	evious 1 Next	96: FROGS Affiliations stat: summary.html 295.0 KB format: html, database: 2	<u>.</u> • / ×
FROGS BIOM to std BIOM Converts a FROGS BIOM in								## Application Software: affiliations_stat.py (version: Command: /usr/local/bioinfo	
<									

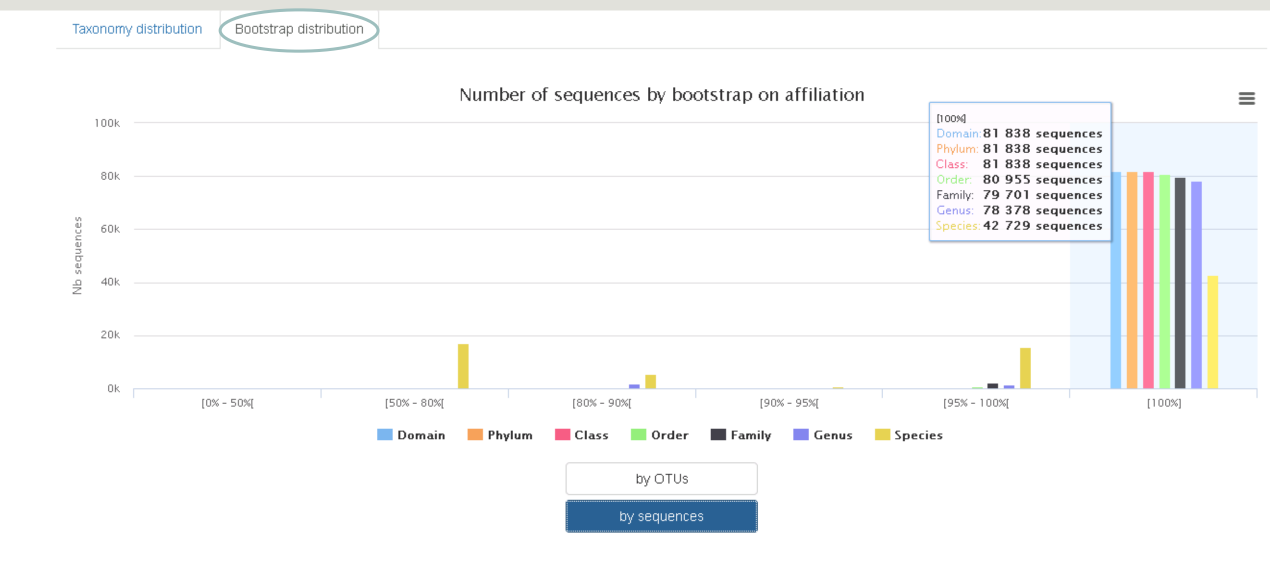
💳 Sigenae - Welcome	e gpascal		Analyze Data Wor	kflow Shared Data <del>-</del>	Visualization - He	elp∓ User∓				Using 88.3 GE
Tools	Taxonomy distributi	on Alignment di	stribution						History	C 0
Split by samples the reads in function of inner barcode.									Formation 9sample 20.3 MB	25
<u>FROGS Pre-process</u> Step 1 in metagenomics analysis: denoising and dereplication.	_		Number of	f OTUs among th	neir alignment re	esults		•	21: FROGS BIOM to TSV: multi hits.tsv	<u>o</u> • 0 %
FROGS Clustering swarm Step 2 in metagenomics	[100%]	0	0	0	0	22	89		20: FROGS BIOM to	<u>o</u> • 0 %
analysis : clustering. FROGS Remove chimera Step	[95% - 100%[	0	0	0	0	20	1	25	TSV: abundance.tsv 19: FROGS Affiliation	
3 in metagenomics analysis : Remove PCR chimera in each sample.	ย [90% - 95%[ ซี	0	0	0	o	10	1	50	stat: summary.html 230.0 KB format: html, databa	1
FROGS Filters Filters OTUs on several criteria.	ی ۱۳۵۵ – ۱۳۵۶ کی ۱۳۵۵ – ۱۳۵۵ کی	0	0	0	0	2	0		## Application Softwaffiliations_stat.py (	ware: version:
FROGS Affiliation OTU Step 4 in metagenomics analysis :	[50% - 80%[	0	0	0	0	0	0	75	1.1.0) Command: /u /bioinfo/src/galaxy-d dist/tools/FROGS/too	dev/galaxy-
Taxonomic affiliation of each OTU's seed by RDPtools and BLAST	[0% - 50%[	0	0	0	0	0	0	100	/affiliations_stat.py /galaxydata/databas /060/dataset_60522	se/files
FROGS BIOM to TSV Converts a BIOM file in TSV file.		[0% - 50%[	[50% - 80%[	[80% – 90%[ Idei	[90% - 95%[ ntity	[95% – 100%[	[100%]	1	output-file /work/g dev/data	galaxy-
FROGS Clusters stat Process some metrics on clusters.				by OTU:	s				HTML file	47 🖻
FROGS Affiliations stat Process some metrics on taxonomies.				by sequen	ces				18: FROGS Affiliati OTU: report.html	ion • 0 X



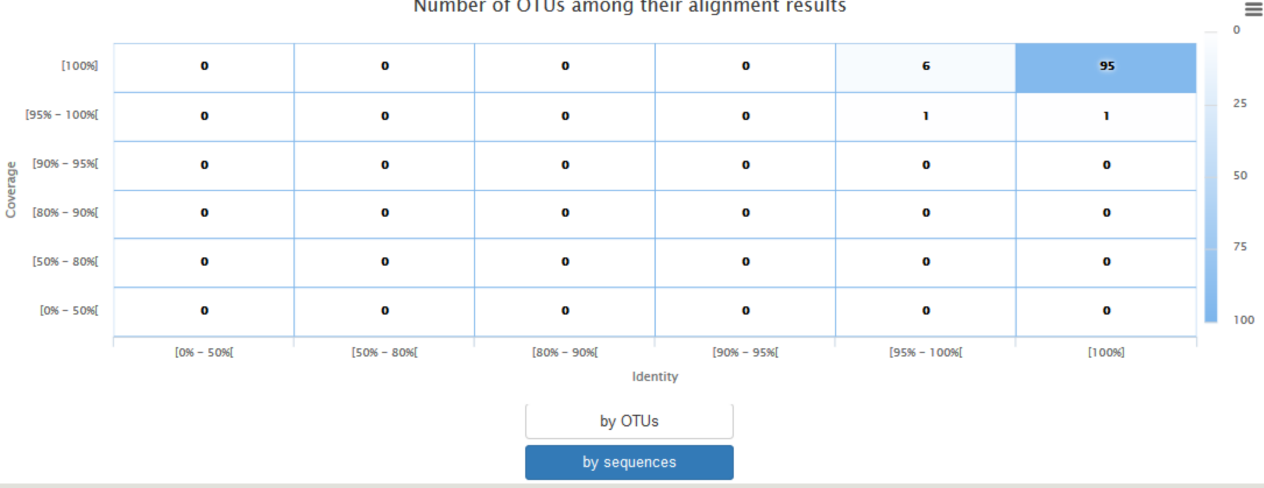






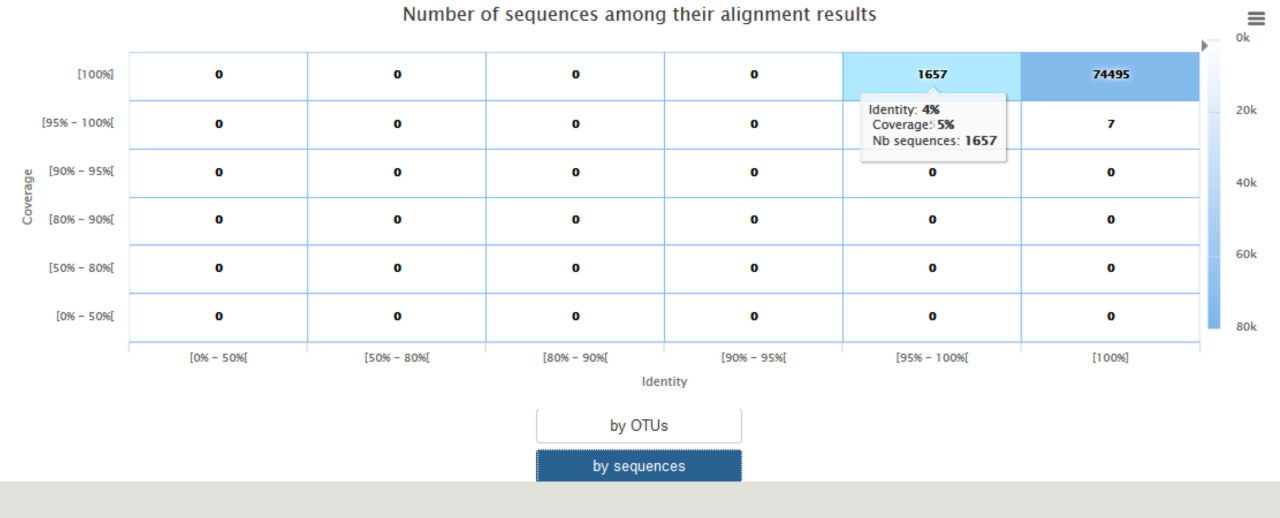






# Number of OTUs among their alignment results

Alignment distributio	nment distributio	t distribution
-----------------------	-------------------	----------------



# TSV to BIOM

FROGS Abundance normalisation 🗶 FROGS Demultiplex reads × Demultiplexing FROGS Affiliations stat 🗶 Seauences file Barcode file Abundance file Select fastq dataset demultiplexed\_archive (data) undemultiplexed archive (data) 🖸 🤇 Normalizat summary (tabular)

#### Upload File from Genotoul × F out1 (bam, txt, tabular, A fastqsanger, csfasta, qual, bed, gff, d gtf, vcf, sam, fasta, pdf, xsq, tar.gz, bw, png, sff, pileup, pileupgz, zip) CO S Data acquisition

0

ROGS Pre-process 🗶	FROGS Clustering swarm
Archive file	Sequences file
dereplicated_file (fasta) 🗆 🧧	Count file
count_file (tabular) 👘 💿 🖓	seed_file (fasta)
summary_file (html) 👘 💿 ( )	abundance_biom (biom1)
	swarms_composition (tabular)
Pre-process	
	Clustering
	Clustering
OM to std BIOM 🗶	Clustering FROGS Clusters stat ¥
OM to std BIOM 🗶	
	FROGS Clusters stat ¥

$\sqrt{\epsilon}$	output_fasta (fasta)
	output_biom (biom1)
ion	summary_file (html)
ustering	swarm 🗶 🗕 F
es file	💳 🔊 s

FROGS Remove chimera	×
Sequences file	
Abundance file	
non_chimera_fasta (fasta)	0
out_abundance_biom (biom1)	0(
out_abundance_count (tabular)	0(
summary_file (html)	0

Chimera

FROGS TSV to BIOM X Abundance TSV File Multi hits TSV File biom\_file (biom1) sequence\_file (fasta) **Convert TSV to Biom** 

Abundance file summary\_file (html)

Affiliation **Statistics** 

FROGS Affiliation OTU OTU seed sequence Abundance file biom\_affiliation (biom1) summary (html)

Affiliation

FROGS Filters × Sequences file Abundance file output\_fasta (fasta) output\_biom (biom1) output\_excluded (tabular) 🖸 output\_summary (html)

**Convert to TSV** 

-multi\_affi\_file (tabular) 🖂 🄇

FROGS BIOM to TSV

Abundance file

Sequences file

tsv\_file (tabular)

FROGS BIO Abundance output bior output\_metadata (tabular) 🖸 **Convert to** 

standard Biom

ers stat 🗙 file le (html) 🛛 Cluster **Statistics** 

196

**Filters** 

# TSV to BIOM

After modifying your abundance TSV file you can again:

- generate rarefaction curve
- sunburst 🔌

Careful :

- <u>do not</u> modify column name
- <u>do not</u> remove column
- take care to choose a taxonomy available in your multi\_hit TSV file
- if deleting line from multi\_hit, take care to not remove a complete cluster without removing all "multi tags" in you abundance TSV file.
- if you want to rename a taxon level (ex : genus "Ruminiclostridium 5;" to genus "Ruminiclostridium;"), do not forget to modify also your multi\_hit TSV file.

# TSV to BIOM

FROGS TSV to BIOM Converts a TSV file in BIOM file. (Galaxy Version 1.0.0)	▼ Options
Abundance TSV File	
29: FROGS BIOM to TSV: abundance.tsv	-
Your FROGS abundance TSV file. Take care to keep intact column name.	
Multi_hits TSV File	
30: FROGS BIOM to TSV: multi_hits.tsv	•
TSV file describinh multi blast hit.	
Extract seed FASTA file	
Yes No	
If there is a 'seed_sequence' column, you can extract seed sequence in a separated FASTA file.	
✓ Execute	

# Your Turn! – 8

PLAY WITH TSV\_TO\_BIOM

# Exercise 8

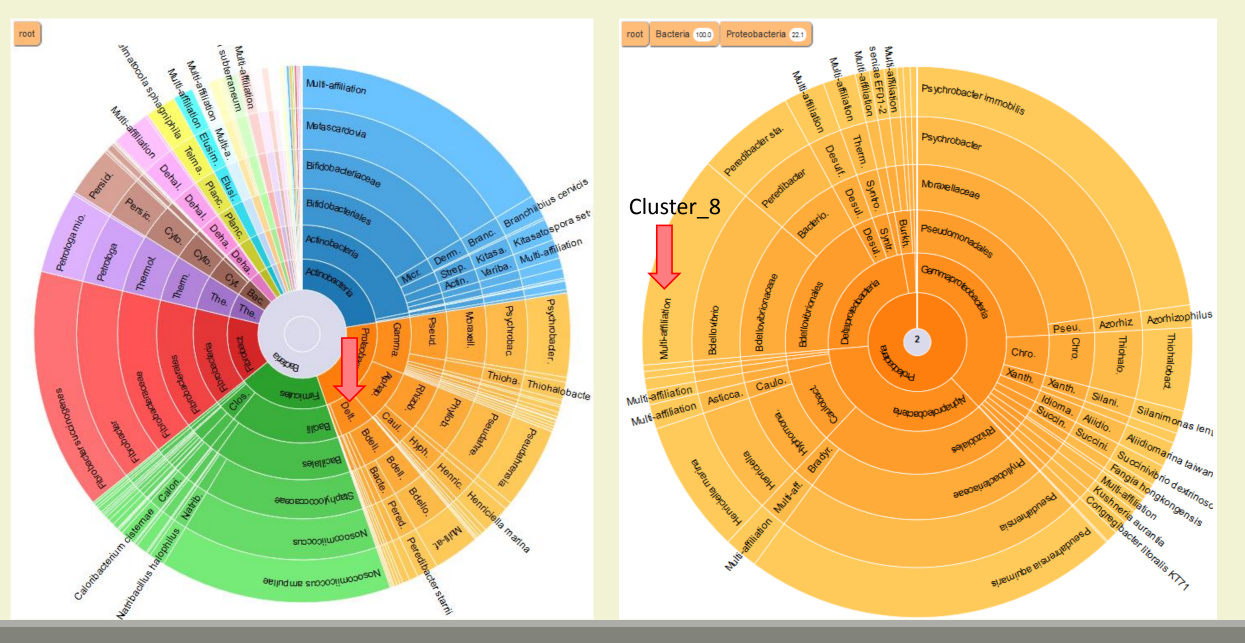
# → objectives : Play with multi-affiliation and TSV\_to\_BIOM

1. Observe in Multi\_hit.tsv and abundance.tsv cluster\_8 annotation

#blast_taxonomy	blast_subject	observation_name	observation_sum
Bacteria; Actinobacteria; Actinobacteria; Bifidobacteriales; Bifidobacteriaceae; Metascardovia; Multi-affiliation	multi-subject	Cluster_1	13337
Bacteria;Fibrobacteres;Fibrobacteria;Fibrobacterales;Fibrobacteraceae;Fibrobacter;Fibrobacter succinogenes	AJ496032.1.1410	Cluster_2	11830
Bacteria;Firmicutes;Bacilli;Bacillales;Staphylococcaceae;Nosocomiicoccus;Nosocomiicoccus ampullae	EU240886.1.1502	Cluster_3	11405
Bacteria;Proteobacteria;Gammaproteobacteria;Pseudomonadales;Moraxellaceae;Psychrobacter;Psychrobacter immobilis	U39399.1.1477	Cluster_4	4125
Bacteria;Thermotogae;Thermotogae;Thermotogales;Thermotogaceae;Petrotoga;Petrotoga miotherma	FR733705.1.1499	Cluster_5	4034
Bacteria; Proteobacteria; Alphaproteobacteria; Rhizobiales; Phyllobacteriaceae; Pseudahrensia; Pseudahrensia aquimaris	GU575117.1.1441	Cluster_6	3966
Bacteria;Bacteroidetes;Cytophagia;Cytophagales;Cytophagaceae;Persicitalea;Persicitalea jodogahamensis	multi-subject	Cluster_7	2433
${\sf Bacteria}; {\sf Proteobacteria}; {\sf Deltaproteobacteria}; {\sf Bdellovibrionales}; {\sf Bdellovibrionaceae}; {\sf Bdellovibrio}; {\sf Multi-affiliation}; {\sf Multi-af$	multi-subject	Cluster_8	2268

Cluster_8	Bacteria; Proteobacteria; Deltaproteobacteria; Bdellovibrionales; Bdellovibrionaceae; Bdellovibrio Bdellovibrio bacteriovorus		CP007656.1036900.1038415	
Cluster_8	Bacteria; Proteobacteria; Deltaproteobacteria; Bdellovibrionales; Bdellovibrionaceae; Bdellovibrio; Bdellovibrio bacteriovorus str. Tiberius		CP002930.1837665.1839157	
Cluster_8	Bacteria; Proteobacteria; Deltaproteobacteria; Bdellovibrionales; Bdellovibrionaceae; Bdellovibrio; Bdellovibrio bacteriovorus str. Tiberius		CP002930.842397.843889	
Cluster_8	Bacteria; Proteobacteria; Deltaproteobacteria; Bdellovibrionales; Bdellovibrionaceae; Bdellovibrio; Bdellovibrio bacteriovorus		AJ292760.1.1334	
Cluster_8	Bacteria; Proteobacteria; Deltaproteobacteria; Bdellovibrionales; Bdellovibrionaceae; Bdellovibrio; Bdellovibrio bacteriovorus		Bdellovibrio bacterio	vorue
Cluster_8	Bacteria; Proteobacteria; Deltaproteobacteria; Bdellovibrionales; Bdellovibrionaceae; Bdellovibrio; Bdellovibrio bacteriovorus	<b></b> /	Buellovibilo bacterio	vorus
Cluster_8	Bacteria; Proteobacteria; Deltaproteobacteria; Bdellovibrionales; Bdellovibrionaceae; Bdellovibrio; Bdellovibrio bacteriovorus		AF084850.1.1436	
Cluster_8	Bacteria; Proteobacteria; Deltaproteobacteria; Bdellovibrionales; Bdellovibrionaceae; Bdellovibrio; Bdellovibrio bacteriovorus HD100		BX842648.123565.125058	
Cluster_8	Bacteria; Proteobacteria; Deltaproteobacteria; Bdellovibrionales; Bdellovibrionaceae; Bdellovibrio; Bdellovibrio bacteriovorus HD100		BX842650.295616.297109	

# 2. Observe le diversity diagramm





# 3. How to change affiliation of cluster 8 ????

# Exercise 8

- 4. Modify multi\_hit.tsv and keep only :
- Cluster\_8 Bacteria;Proteobacteria;Deltaproteobacteria;Bdellovibrionales;Bdellovibrionaceae;Bdellovibrio;Bdellovibrio bacteriovorus CP007656.1036900.1038415

Careful, <u>no quotes</u> around text !!!

- 5. Upload the new multihit file.
- 6. Create a new biom with a TSV\_to\_BIOM tool
- 7. Launch again the affilation\_stat tool on this new biom
- 8. Observe the diversity diagram



# Normalization

FROGS Demultiplex reads FF × Demultiplexing Se Barcode file Select fastq dataset Ab demultiplexed\_archive (data) οι undemultiplexed archive (data) 🖸 🤇 οι Normalization summary (tabular) su

×

ROGS Abundance normalisatio	in 🕽
equences file	
oundance file	
utput_fasta (fasta)	
utput_biom (biom1)	
ummary_file (html)	E

×

00

#### Upload File from Genotoul × out1 (bam, txt, tabular, fastqsanger, csfasta, qual, bed gtf, vcf, sam, fasta, pdf, xsg, tar bw, png, sff, pileup, pileupgz, zi

#### Data acquisition

FROGS BIOM to TSV

-multi\_affi\_file (tabular) 🖂 🤇

**Convert to TSV** 

0

Abundance file

Sequences file

tsv\_file (tabular)

	) Archive file	
, gff, 🔁	dereplicated_file (fast	a) 🖸
ip)	count_file (tabular)	
	summary_file (html)	0
	Pre-process	
FROGS E	BIOM to std BIOM 🗙	
🕑 Abunda	nce file	
output_	biom (biom1) 🛛 🖸 🖸	
output_	metadata (tabular) 🗇 🤇	

FROGS Pre-process

**Convert to** standard Biom

	FROGS Clusters stat 🗙
8	Abundance file
Ħ	summary_file (html) 🜼
	Cluster Statistics

FROGS Clustering swarm

abundance\_biom (biom1)

swarms\_composition (tabular)

Sequences file

seed\_file (fasta)

Clustering

Count file

## FROGS Remove chimera × Sequences file Abundance file non\_chimera\_fasta (fasta) out\_abundance\_biom(biom1) out\_abundance\_count (tabular) 🗇 🤇 summary\_file (html)

## Chimera

FROGS TSV to BIOM X Abundance TSV File Multi hits TSV File biom\_file (biom1) sequence\_file (fasta) **Convert TSV to Biom** 

#### FROGS Affiliations stat 🗶

Abundance file summary\_file (html)

## Affiliation **Statistics**

FROGS Affiliation OTU OTU seed sequence Abundance file biom\_affiliation (biom1) summary (html)

Affiliation

FROGS Filters × Sequences file Abundance file output\_fasta (fasta) output\_biom (biom1) output\_excluded (tabular) 🖸

output\_summary (html)

**Filters** 

# Normalization

Conserve a predefined number of sequence per sample:

- update Biom abundance file
- update seed fasta file

May be used when :

- Low sequencing sample
- Required for some statistical methods to compare the samples in pairs

# Your Turn! – 9

LAUNCH NORMALIZATION TOOL

# Exercise 9

Launch Normalization Tool

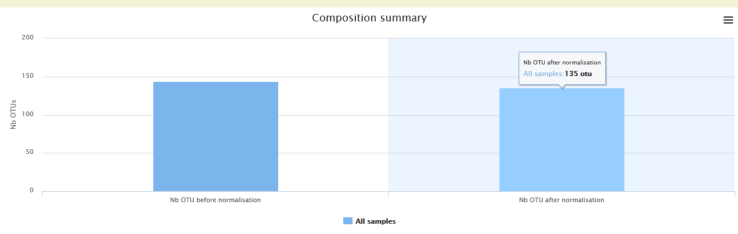
- 1. What is the smallest sequenced samples ?
- 2. Normalize your data from Affiliation based on this number of sequence
- 3. Explore the report HTML result.
- 4. Try other threshold and explore the report HTML result What do you remark ?

FROGS Abundance normalisation (Galaxy Version 1.1.1)	▼ Options
Sequences file	
17: FROGS Filters: sequences.fasta	•
Sequences file to normalize (format: fasta).	
Abundance file	
22: FROGS Affiliation OTU: affiliation.biom	-
Abundances file to normalize (format: BIOM).	
Number of reads	
9088	
The final number of reads per sample.	
✓ Execute	

FROGS Abundance normalisation (Galaxy Version 1.1.1)	✓ Options
Sequences file	
17: FROGS Filters: sequences.fasta	•
Sequences file to normalize (format: fasta).	
Abundance file	
22: FROGS Affiliation OTU: affiliation.biom	-
Abundances file to normalize (format: BIOM).	
Number of reads	
2000	
The final number and s per sample.	
✓ Execute	
Or, this number can be chosen according to the rarefaction	
curve. For example, we can choose the smallest number of	
sequences that still retain all the genus.	
sequences that still retain all the genus.	

2	1	1
2	-	ь.

0	Nb OTU before normalisation			Nb OTU after normalisation			
		All samp	es				
							acsv
Show 10 - entries				Search	:	2	≅CSV
Composition by sample							
Sample	<b>▲</b> 1	Nb OTU before normalisatio	n	Nb OTU after normalisation	ı		÷
100_10000seq_sampleA1_cutadapt	1	144		135			
100_10000seq_sampleA2_cutadapt	1	144		135			
100_10000seq_sampleA3_cutadapt	1	144		135			
100_10000seq_sampleB1_cutadapt	1	144		135			
100_10000seq_sampleB2_cutadapt	1	144		135			
100_10000seq_sampleB3_cutadapt	1	144		135			
100_10000seq_sampleC1_cutadapt	1	144		135			
100_10000seq_sampleC2_cutadapt	1	144		135			
100_10000seq_sampleC3_cutadapt	1	144		135			
Showing 1 to 9 of 9 entries					Previous	1	Next



# Filters on affiliations

Do not forget, with filter tool we can filter the data based on their affiliation

FROGS Filters Filters OTUs on several criteria. (Galaxy Version 1.2.0)	- Options
Sequences file	
9: FROGS Remove chimera: non_chimera.fasta The sequence file to filter (format: fasta).	
ibundance file	
	-
10: FROGS Remove chimera: non_chimera_abundance.biom	
he abundance file to filter (format: BIOM).	
** THE FILTERS ON OTUS IN SAMPLES, OTUS SIZE and SEQUENCE PERCENTAGE	
Apply filters	Abundance filters
you want to filter OTUs on their abundance and occurrence.	
Minimum number of samples	
Fill the field only if you want this treatment. Keep OTU present in at least this number of samples.	
Minimum proportion/number of sequences to keep OTU	
Fill the field only if you want this treatment. Use decimal notation for proportion (example: 0.01 for keep OTU with at least 1% o Use integer notation for number of sequence (example: 2 for keep OTU with at least 2 sequences, so remove single singleton).	of all sequences) ;
N biggest OTU	
Fill the fields only if you want this treatment. Keep the N biggest OTU.	
•• THE FILTERS ON RDP	
Apply filters	RDP affiliation filters
you want to filter OTUs on their taxonomic affiliation produced by RDP.	Ref annation meets
Rank with the bootstrap filter	
Nothing selected	•
Minimum bootstrap % (between 0 and 1)	
** THE FILTERS ON BLAST	
Apply filters ' you want to filter OTUs on their taxonomic affiliation produced by Blast.	<b>BLAST</b> affiliation filters
Maximum e-value (between 0 and 1)	
Fill the field only if you want this treatment	
Minimum identity % (between 0 and 1)	
Fill the field only if you want this treatment	
Minimum coverage % (between 0 and 1)	
Fill the field only if you want this treatment	
Minimum alignment length	
Fill the field only if you want this treatment	
** THE FILTERS ON CONTAMINATIONS	
Apply filters	Contamination filter
	Contamination Inter
	Containination miter
you want to filter OTUs on classical contaminations.	
f you want to filter OTUs on classical contaminations. Cotaminant databank	

# Exercise 10

- 1. Apply filters to keep only data with perfect alignment.
- 2. How many clusters have you keep?

	ilters OTUs on several criteria. (Galaxy Version 1.2.0)	
equences file		
C 2 C	17: FROGS Filters: sequences.fasta	-
he sequence file	e to filter (format: fasta).	
bundance file		
C 2 C	22: FROGS Affiliation OTU: affiliation.biom	•
he abundance f	file to filter (format: BIOM).	
** THE FILTER	RS ON OTUS IN SAMPLES, OTUS SIZE and SEQUENCE PERCENTAGE	
No filters		•
f you want to filt	ter OTUs on their abundance and occurrence.	
** THE FILTER	RS ON RDP	
No filters		
f you want to filt	ter OTUs on their taxonomic affiliation produced by RDP.	
f you want to filt *** THE FILTER		
** THE FILTER		
*** THE FILTER Apply filters f you want to filt	RS ON BLAST	
Apply filters f you want to filt Maximum e-va	RS ON BLAST ter OTUs on their taxonomic affiliation produced by Blast. alue (between 0 and 1)	•
*** THE FILTER Apply filters f you want to filt Maximum e-va Fill the field on	RS ON BLAST ter OTUs on their taxonomic affiliation produced by Blast. alue (between 0 and 1)	
*** THE FILTER Apply filters f you want to filt Maximum e-va Fill the field on	RS ON BLAST ter OTUs on their taxonomic affiliation produced by Blast. alue (between 0 and 1)	
Apply filters f you want to filt Maximum e-va Fill the field on Minimum ident	AS ON BLAST ter OTUs on their taxonomic affiliation produced by Blast. alue (between 0 and 1) Ily if you want this treatment tity % (between 0 and 1)	· · · · · · · · · · · · · · · · · · ·
Apply filters f you want to filt Maximum e-va Fill the field on Minimum ident	RS ON BLAST ter OTUs on their taxonomic affiliation produced by Blast. alue (between 0 and 1)	· · · · · · · · · · · · · · · · · · ·
Apply filters f you want to filt Maximum e-va Fill the field on Minimum ident 1 Fill the field on	AS ON BLAST ter OTUs on their taxonomic affiliation produced by Blast. alue (between 0 and 1) Ily if you want this treatment tity % (between 0 and 1)	
Apply filters f you want to filt Maximum e-va Fill the field on Minimum ident 1 Fill the field on	AS ON BLAST ter OTUs on their taxonomic affiliation produced by Blast. alue (between 0 and 1) Ily if you want this treatment tity % (between 0 and 1) Ily if you want this treatment	
** THE FILTER Apply filters f you want to filt Maximum e-va Fill the field on Minimum ident 1 Fill the field on Minimum cove	AS ON BLAST ter OTUs on their taxonomic affiliation produced by Blast. alue (between 0 and 1) Ily if you want this treatment tity % (between 0 and 1) Ily if you want this treatment	

Fill the field only if you want this treatment

# Tool descriptions



### What it does

FROGS Pre-process filters and dereplicates amplicons for use in diversity analysis.

### Inputs/Outputs

#### Inputs

By sample your sequences and their qualities.

#### Illumina inputs

Usage: The amplicons have been sequenced in paired-end. The amplicon expected length is inferior than the R1 and R2 length. R1 and R2 can be merge by the common region.
 Files: One R1 and R2 by sample (format <u>FASTQ</u>)
 Example: splA\_R1.fastq.gz, splA\_R2.fastq.gz, splB\_R1.fastq.gz, splB\_R2.fastq.gz

#### OR

 Usage:
 The single end sequencing cover all the amplicons or the R1 and R2 have already been overlaped.

 Files:
 One sequence file by sample (format FASTQ).

Example: splA.fastq.gz, splB.fastq.gz

#### 454 inputs

Files: One sequence file by sample (format <u>FASTQ</u>) Example: splA.fastq.gz, splB.fastq.gz

These files must be added sample by sample or provide in an archive file (tar.gz). Remark: In an archive if you use R1 and R2 files they names must end with \_R1 and \_R2.

#### Outputs

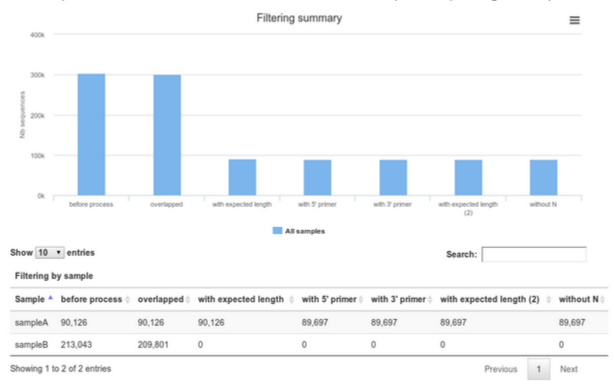
Sequence file (dereplicated.fasta):

Only one file with all samples sequences (format <u>FASTA</u>). These sequences are dereplicated: strictly identical sequence are represented only one and the initial count is kept in count file.

Count file (count.tsv):

This file contains the count of all uniq sequences in each sample (format TSV).

Summary file (excluded\_data.html):



This file presents the ordered filters and the number of sequences passing these (format HTML).

## <sup>1</sup> How it works

Steps	Illumina	454
1	For uncontiged data: contig read1 and read2 with a maximum of 10% mismatch in the overlaped region ( <u>FLASh</u> )	/
2	Filter contig sequence on its length which must be between Minimum amplicon size" and "Maximum amplicon size"	/
3	Remove sequences where the two primers are not persent and remove primers sequence ( <u>cutadapt</u> ). The primer search accept 10% of differences	Remove sequence where the two primers are not persent, remove primers sequence and reverse complement the sequences with strand - ( <u>cutadapt</u> ). The primer search accept 10% of differences
4	Filter sequences on its length and with ambiguous nucleotids	filter sequences on its length, with ambiguous nucleotids, with at least one homopolymer with size >7nt and with distance between two poor qualities ()< 10) of <= 10 nt
5	Dereplicate sequences	Dereplicate sequences

### <sup>1</sup> Advices/details on parameters

#### Primers parameters

The primers must provided in 5' to 3' orientation.

Example:

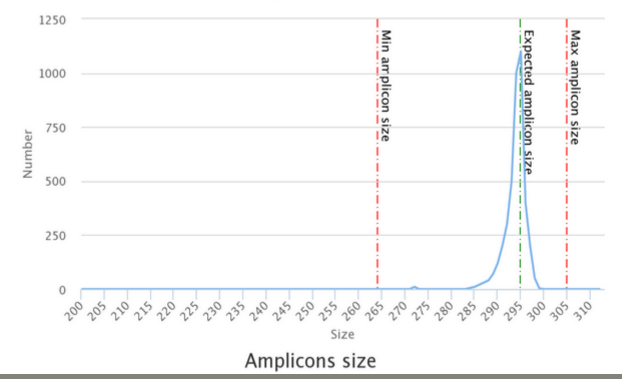
5' ATGCCC GTCGTCGTAAAATGC ATTTCAG 3'

Value for parameter 5' primer: ATGCC

Value for parameter 3' primer: ATTTCAG

### Amplicons sizes parameters

The two following images shown two examples of perfect values fors sizes parameters.



Amplicons size

# Workflow creation

Workflow Canvas | frogs v1.0

#### Details

				Tool: (beta) FROGS Filters (beta)
Upload File <b>X</b> Out1 (bam, txt, fastqsanger, Status and the state of t	ocess X	(beta) FROGS Clustering swarm X (beta) Sequences file	(beta) FROGS Clusters stat (beta) Cluster file summary_file (html)	Version: 1.0.0 None:  Biom File Data input 'biom' (txt) Fasta File
csfasta, qual, bed, gff, gtf, vcf, sam, fasta, pdf, xsq, tar.gz, bw, png) count_file (tabular) summary_file (html)		abundance_biom (txt) seed_file (fasta) swarms_composition (tabular)	(beta) FROGS Remove chimera ★ (beta) Sequences file Abundance file non_chimera_fasta (fasta)	Data input 'fasta' (fasta) Remove phiX:  PhiX databank:  phiX  *** THE FILTERS ON OTUS IN SAMPLES, OTUS SIZE and
	fasta_output (fasta)       web (html)	(beta) ¥ (beta) FROGS Affilia (beta) OTU abondance in OTU seed sequence	biom format	SEQUENCE PERCENTAGE : Apply filters Remove OTUs that are not present at least in XX samples; how many samples do you choose? : When sorted by abundance, how many OTU do you want to keep ?:
	(beta) Cluster	FROGS Clusters stat	:) summary_file (html)	proportion/number of sequences threshold to remove an OTU: ▼ 0.00005 *** THE FILTERS ON RDP : No filters ▼ *** THE FILTERS ON BLAST : No filters ▼

# Your Turn! – 11

CREATE YOUR OWN WORKFLOW !

Exercise 11					
			2		
- Galaxy Sigenae - Welcome gnas	Cal Analyza Data Worldlow Charod Data = View	alization - Hole - Hoor-		EEE Ulaina	10.2.0
<b>g</b> Galaxy Sigenae - Welcome gpas Your workflows	Ca Analyze Data Workflow Shared Data - Visu	alization∓ Help∓ User∓	Create new workflow	Using	
	Cal Analyze Data Workflow Shared Data - Visu	alization∓ Help∓ User∓	© Create new workflow # of Steps		18.3 GE
/our workflows	Ca Analyze Data Workflow Shared Data - Visu	alization			
Your workflows	Cal Analyze Data Workflow Shared Data - Visu	alization	# of Steps		

No workflows have been shared with you.

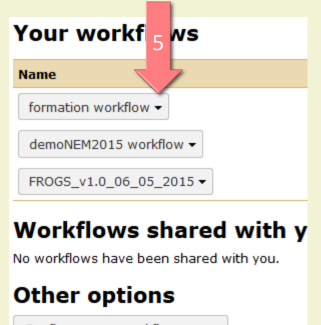
### **Other options**

Configure your workflow menu

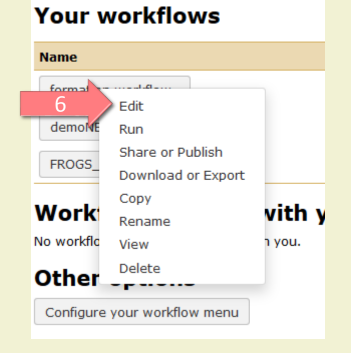
# Exercise 11



# Exercise 11



Configure your workflow menu



#### Upload File from Genotoul

out1 (bam, txt, tabular, fastqsanger, csfasta, qual, bed, gff, gtf, vcf, sam, fasta, pdf, xsq, tar.gz, bw, png, sff, pileup, pileupgz, zip) FROGS Pre-process

×

dereplicated\_file (fasta) 0 0 count\_file (tabular) 0 0

summary\_file (html)

FROGS Clustering swarm

Count file

×

💙 seed\_file (fasta)

abundance\_biom (biom1) O

×

 FROGS Remove chimera
 X

 Sequences file
 Abundance file

 non\_chimera\_fasta (fasta)
 O

 out\_abundance\_biom (biom1)
 O

 out\_abundance\_count (tabular)
 O

 summary\_file (html)
 O

FROGS Affiliation OTU X OTU seed sequence Abundance file biom\_affiliation (biom1) summary (html)

>



×

#### Upload File from Genotoul

out1 (bam, txt, tabular, fastqsanger, csfasta, qual, bed, gff, gtf, vcf, sam, fasta, pdf, xsq, tar.gz, bw, png, sff, pileup, pileupgz, zip)

×

 FROGS Pre-process
 X

 Archive file

 dereplicated\_file (fasta)

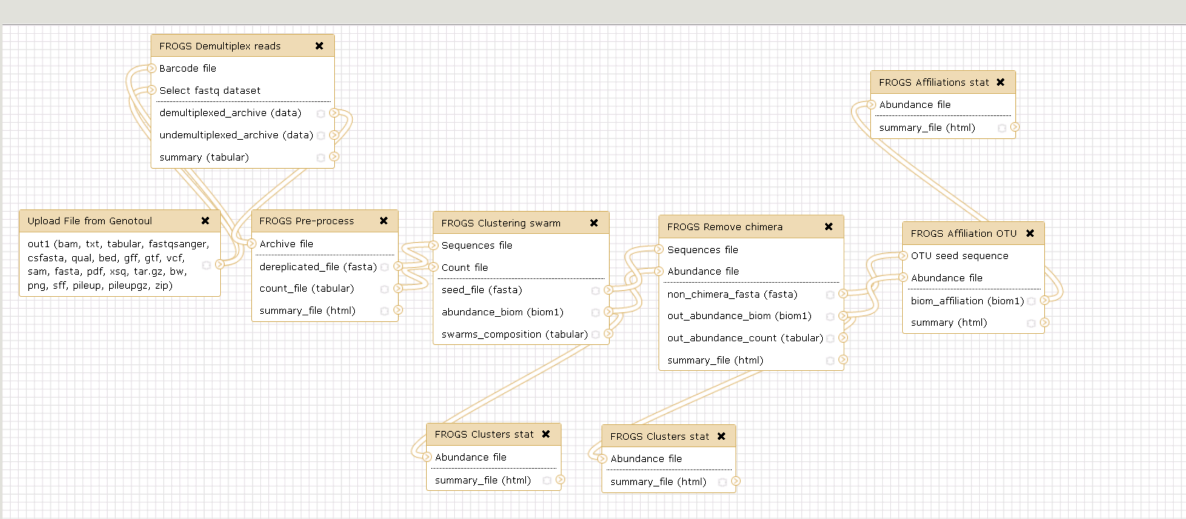
 count\_file (tabular)

 summary\_file (html)

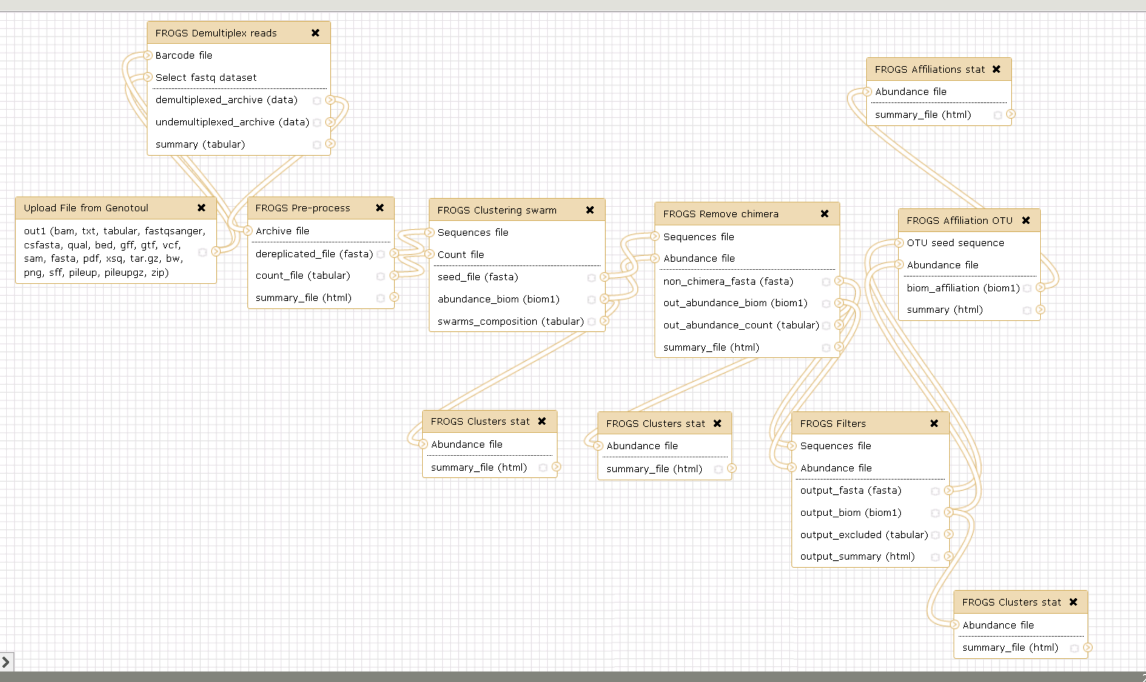
FROGS Clustering swarm Sequences file Count file seed\_file (fasta) abundance\_biom (biom1)

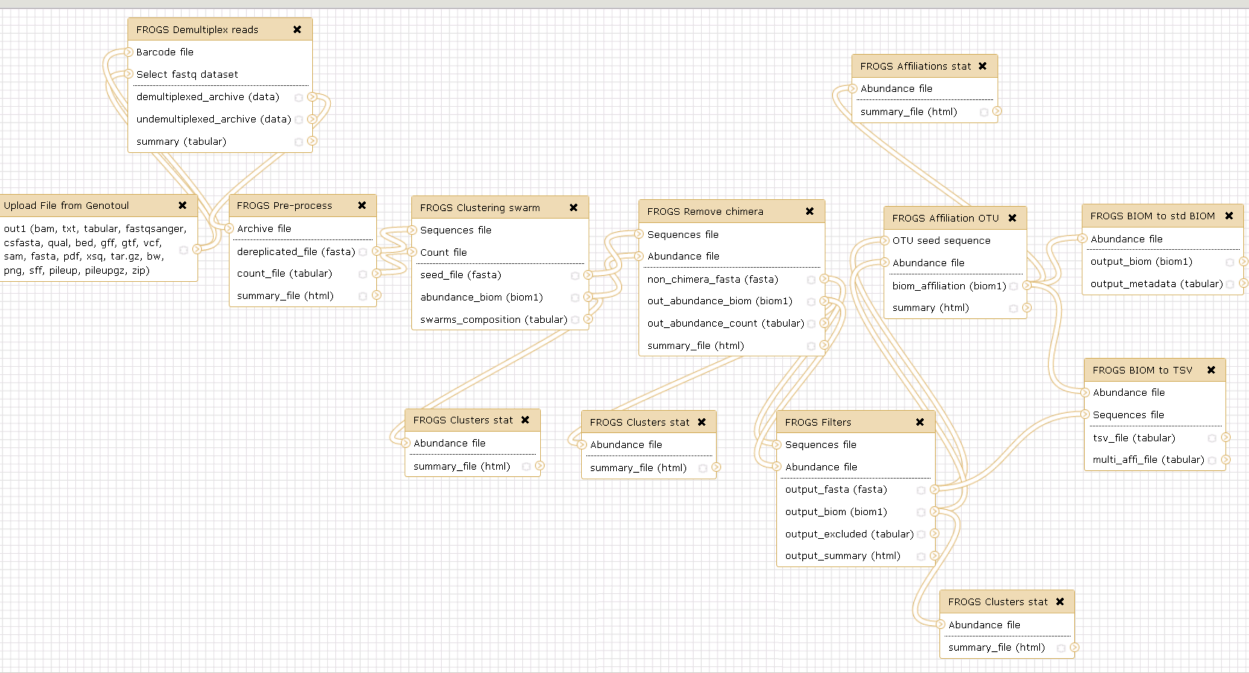
swarms\_composition (tabular) 🗆 📀

	FROGS Remove chimera	×		FROGS Affiliation OTU 🗙
Ç	Sequences file			OTU seed sequence
-0	Abundance file		8	Abundance file
	non_chimera_fasta (fasta)	00	={{	biom_affiliation (biom1) 🖸
	out_abundance_biom (biom1)	00		summary (html)
	out_abundance_count (tabular	) 🖸 🤇		
	summary_file (html)	00	<b>)</b>	

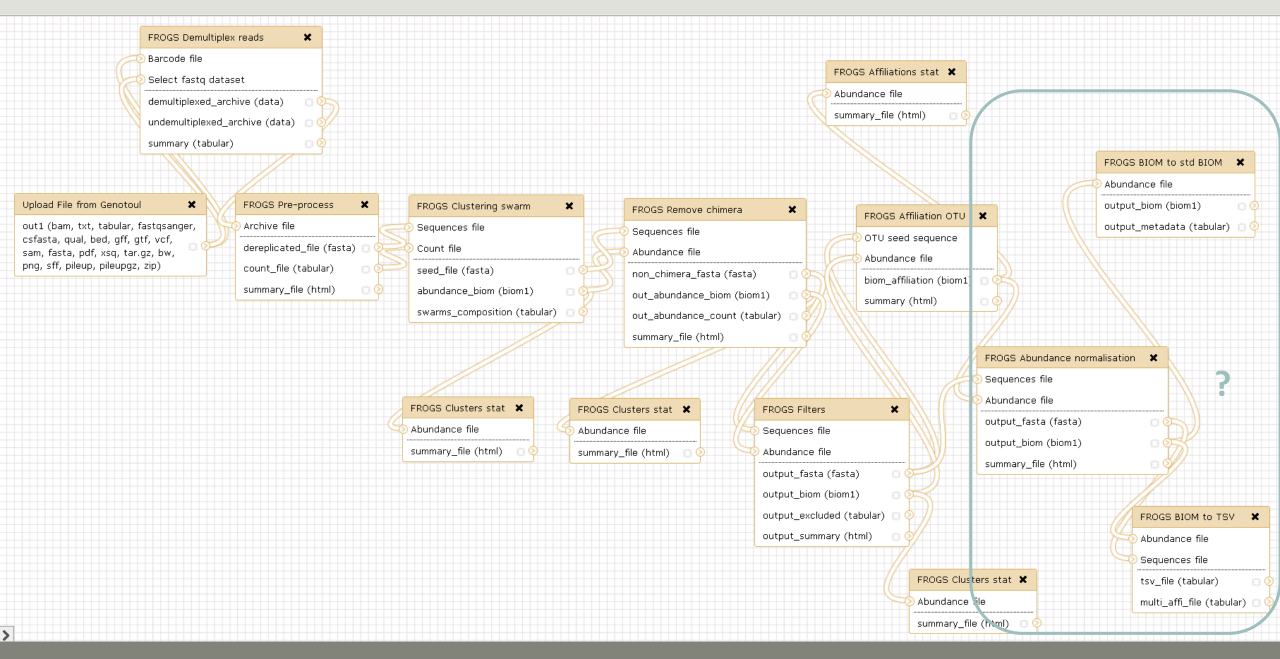


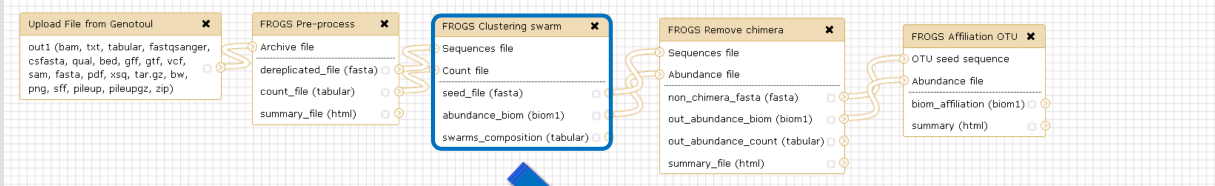






>





• Fixe parameter ?

FROGS Clustering swarmStep 2 in metagenomicsanalysis : clustering. (GalaxyVersion 2.3.0)

#### Sequences file Data input 'sequence\_file' (fasta) The sequences file (format: fasta).

Count file

Data input 'count\_file' (tabular) It contains the count by sample for each sequence (format: TSV).

#### Aggregation distance

#### Set at Runtime

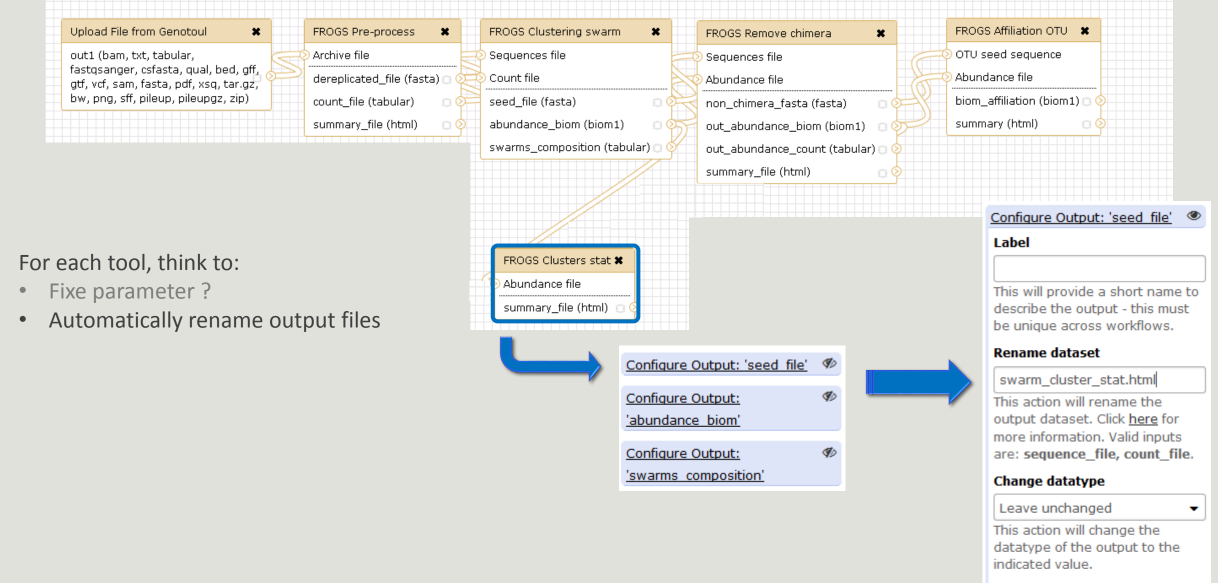
?

Maximum number of differences between sequences in each aggregation step.

Performe denoising clustering step?

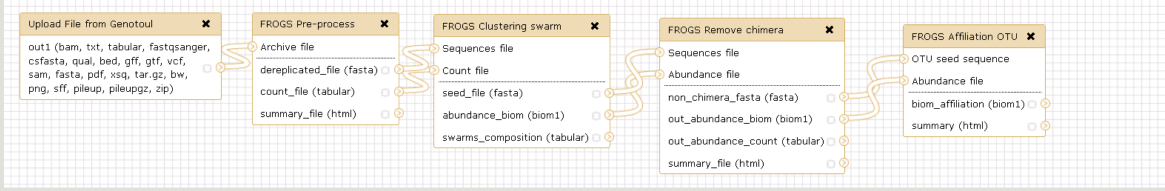
Yes No

If checked, clustering will be perform in two steps, first with

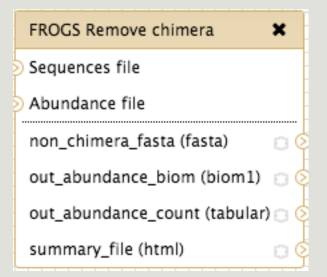


#### Tags

This action will set tags for the dataset.

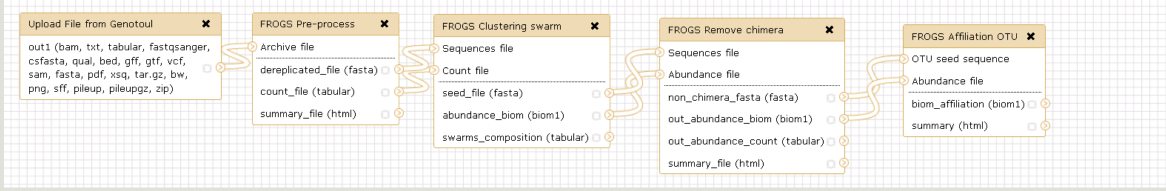


- Fixe parameter ?
- Automatically rename output files
- Hide intermediate files ?



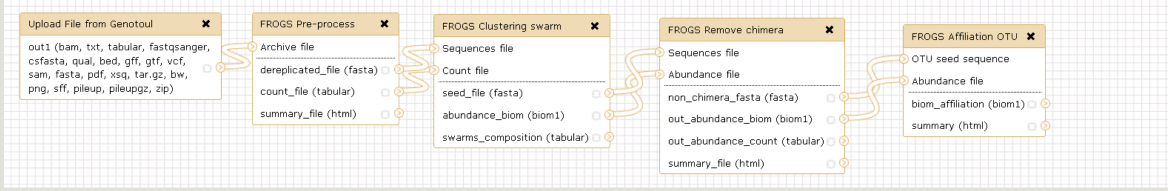


11: FROGS Remove chimera: report.html	• / %
10: FROGS Remove chimera: non chimera abundance.biom	• / %
<u>9: FROGS Remove chimera:</u> non_chimera.fasta	• / ×

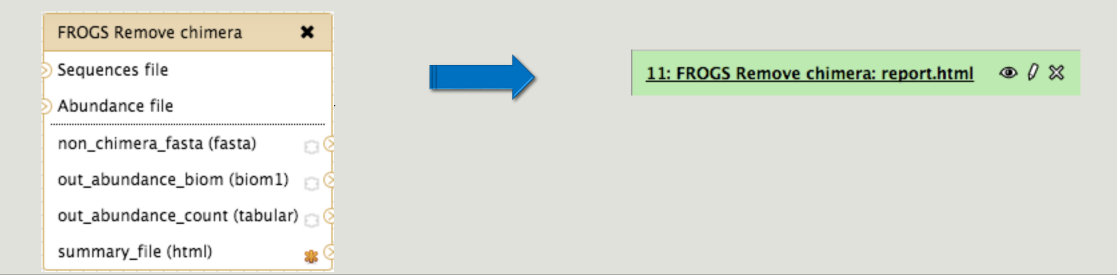


- Fixe parameter ?
- Automatically rename output files
- Hide intermediate files ?

FROGS Remove chimera
Sequences file
> Abundance file
non_chimera_fasta (fasta) 🛛 💿 🔯
out_abundance_bio_Mark_dataset as a workflow output. All unmarked datasets
out_abundance_count (tabwill be hidden.
summary_file (html)



- Fixe parameter ?
- Automatically rename output files
- Hide intermediate files ?



# Download your data

## You have to download one per one your files

	55: FROGS Affiliation @ 0 🛛
	OTU:
	excluded data report.html
	11.4 KB
	format: html, database: ?
	## Application Software:
	affiliation_OTU.py (version: 0.4.0)
	Command: /usr/local/bioinfo
	/src/galaxy-test/galaxy-dist/tools
	/FROGS/affiliation_OTU.py
	reference /save/galaxy-
	test/bank/FROGS/silva_119-1
	/prokaryotes
	/silva_119-1_prokaryotes.fasta
	abundance
	, 🖬 🛈 🧶 📄
2	
	HTML file

# FROGS BIOM to Standard BIOM

# FROGS biom to standard Biom

## This step is required to run R

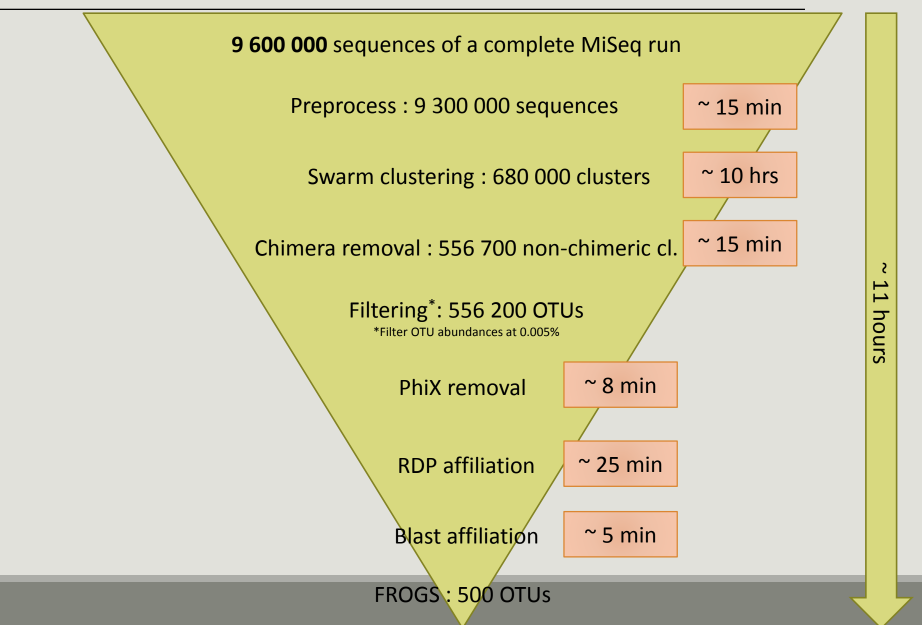
ROGS BIOM to std BIOM Converts a FROGS BIOM in fully compatible BIOM. (Galaxy Version 1.1.0)
Abundance file
22: FROGS Affiliation OTU: affiliation.biom
The FROGS BIOM file to convert (format: BIOM).
✓ Execute
<u>43: FRO</u>
blast m
42: FRO

# Some figures

# Some figures - Fast

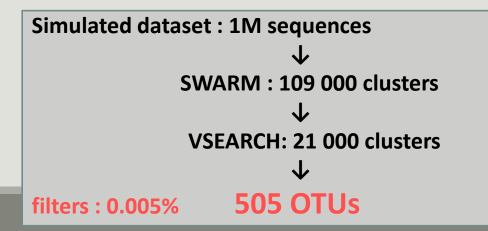
NB SEQ	TIME with complete pipeline without Filters
50 000	40 min
400 000	4 hrs
3 500 000	2 days
10 000 000	5 days

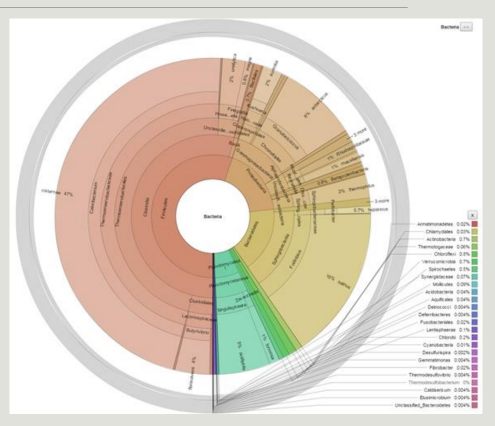
# Speed on real datasets



# Simulated datasets, for testing FROGS' Accuracy

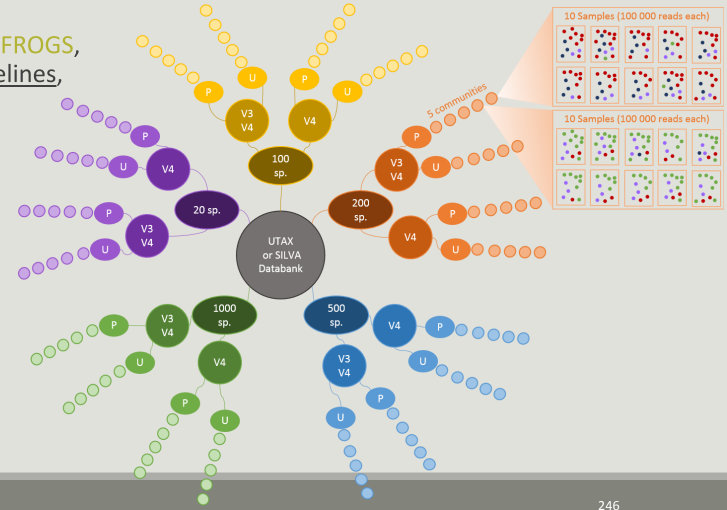
- 500 species, covering all bacterial phyla
- Power Law distribution of the species abundances
- Error rate calibrated with real sequencing runs
- 20% chimeras
- 10 samples of 100 000 sequences each (IM sequences)





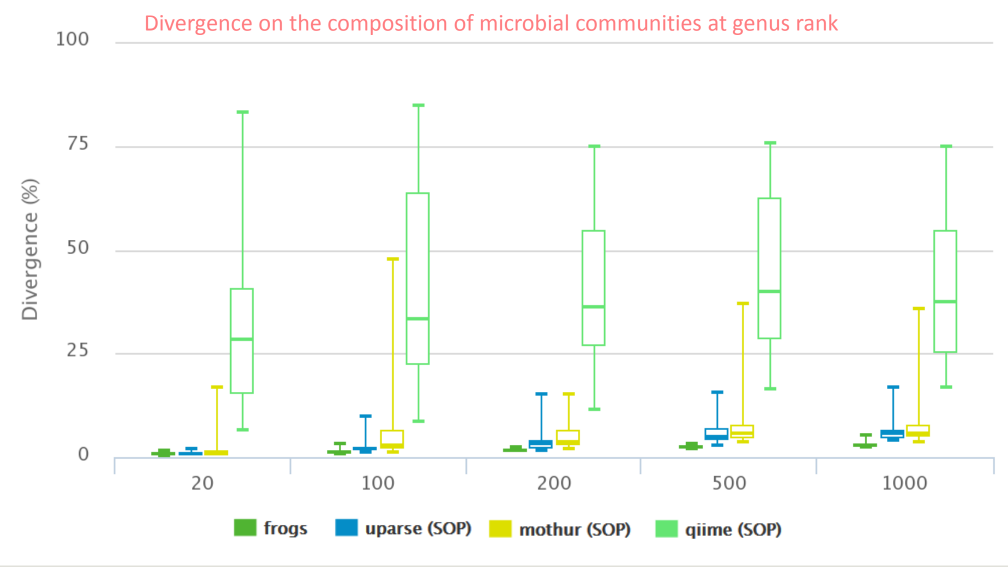
# FROGS' Accuracy

- 1.10<sup>+8</sup> synthetic sequences were treated with FROGS, UPARSE and MOTHUR, QIIME, with their guidelines, to compare their performances
- 20, 100, 200, 500 or 1000 different species
- power law or a uniform distribution
- 5 to 20% of chimera
- $\rightarrow$  Divergence on the composition of microbial communities at the different taxonomic ranks



#### V3V4 Power Law

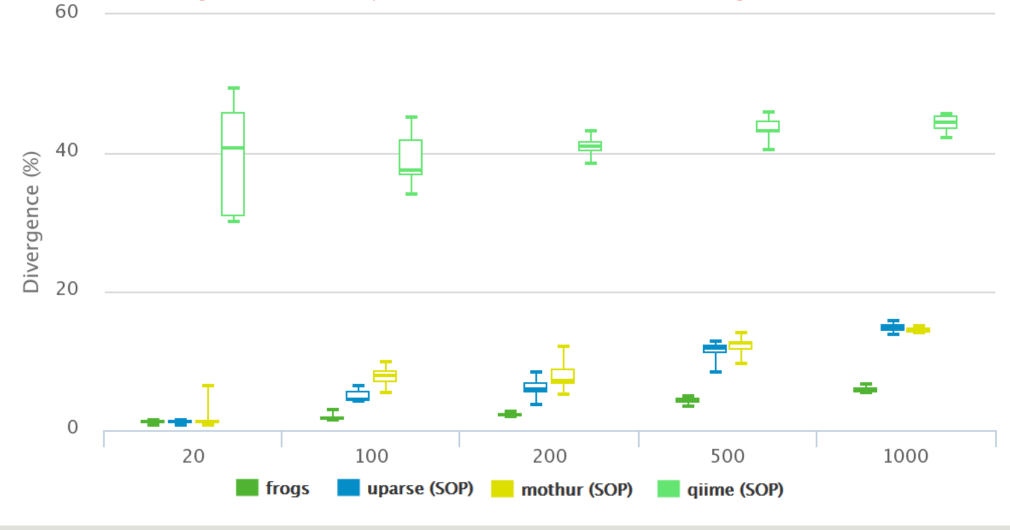
## Affiliations divergence



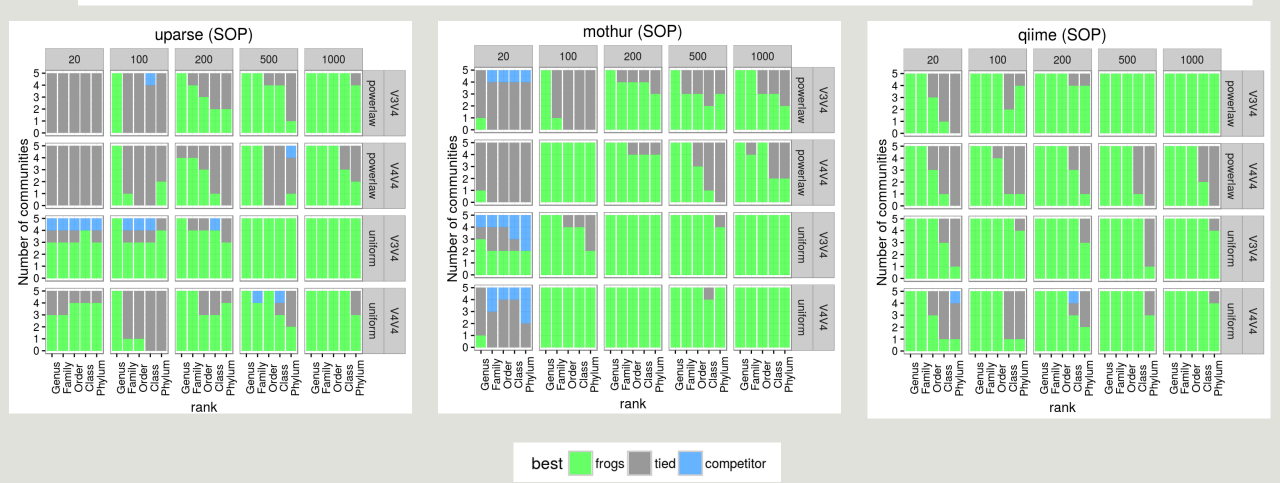
## Affiliations divergence

V3V4 Uniform

Divergence on the composition of microbial communities at genus rank



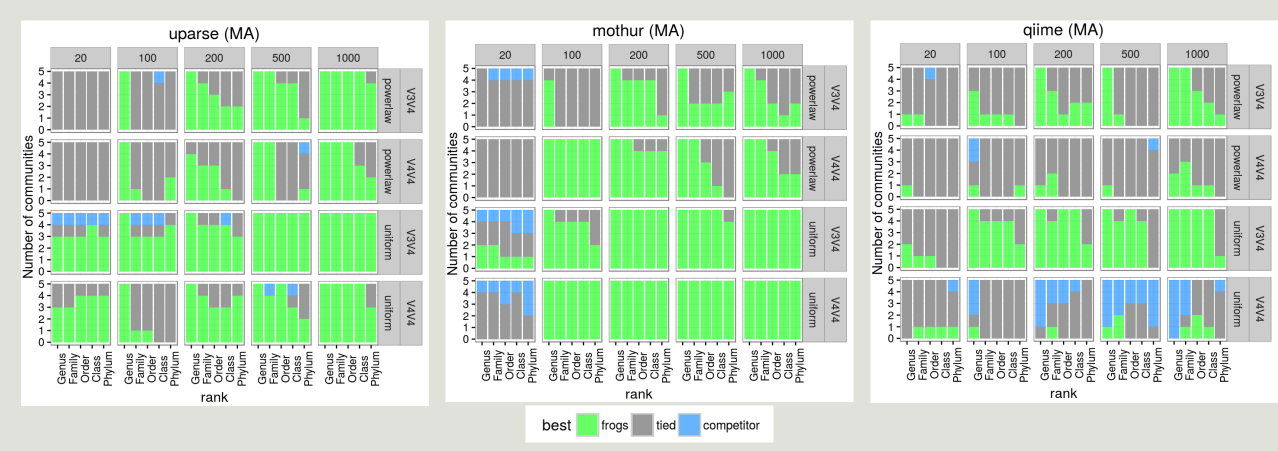
### The results of non-parametric paired tests (signed rank test) of Affiliation divergence on simulated data from UTAX



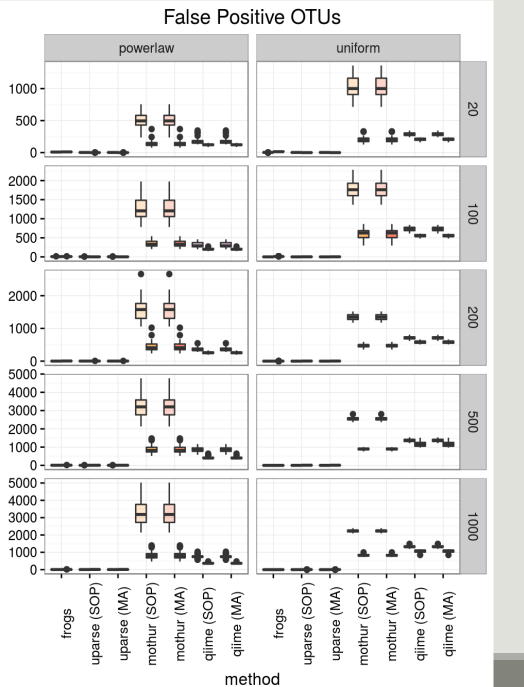
FROGS performs as well as or better than UPARSE, MOTHUR and QIIME in most settings. The only condition in which FROGS does worse than UPARSE and MOTHUR is small community size (20).

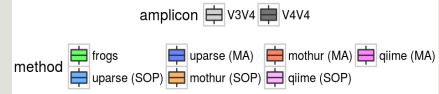
### The results of non-parametric paired tests (signed rank test) of Affiliation divergence on simulated data from UTAX

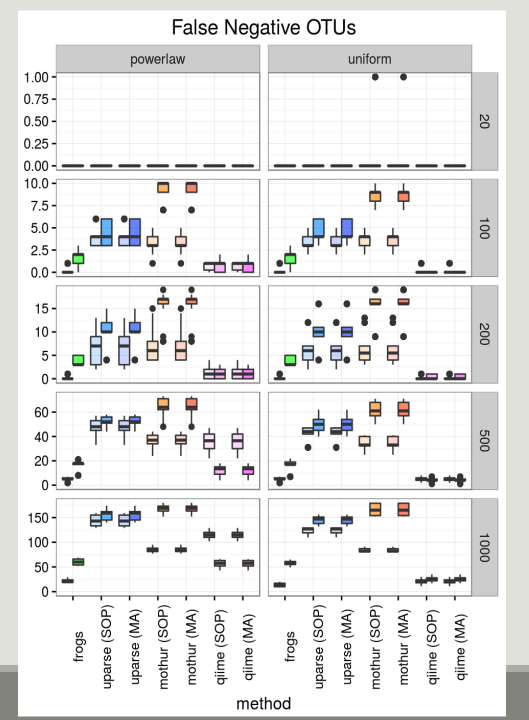
### With FROGS multi-Affiliation



QIIME (MA) with large communities (size > 200) with uniform abundance using the V4 region is better than FROGS. The differences, although significant, are small in that case: less than 2 percentage points in all cases and most marked at the Genus level where the divergences of both FROGS and QIIME (MA) are already quite moderate (6~10%).









# Conclusions



# Why Use FROGS ?

- User-friendly
- Fast
- 454 data and Illumina data
  - sequencing methods change but same tool
  - easier for comparisons
- Clustering without global threshold and independent of sequence order
- New chimera removal method (Vsearch + cross-validation)

- Filters tool
- Multi-affiliation with 2 taxonomy affiliation procedures
- Cluster Stat and Affiliation Stat tools
- A lot of graphics
- Independant tools



# How to cite FROGS

In waiting for the publication:

Pipeline FROGS on <a href="http://sigenae-workbench.toulouse.inra.fr/">http://sigenae-workbench.toulouse.inra.fr/</a>

Github: <u>https://github.com/geraldinepascal/FROGS.git</u>

Poster FROGS: Escudie F., Auer L., Bernard M., Cauquil L., Vidal K., Maman S., Mariadassou M., Combes S., Hernadez-Raquet G., Pascal G., 2016. FROGS: Find Rapidly OTU with Galaxy Solution. In: ISME-2016 Montreal, CANADA,

http://bioinfo.genotoul.fr/wp-content/uploads/FROGS\_ISME2016\_poster.pdf



## To contact

FROGS:

frogs@toulouse.inra.fr

Galaxy:

sigenae-support@listes.inra.fr

Newsletter – demande d'abonnement:

mailto:sympa@listes.inra.fr?subject=sub%20frogs-newsletter

frogs-newsletter-request@listes.inra.fr



# Next training sessions

3<sup>rd</sup> to 6<sup>th</sup> July 2017 4 days

0.5 Galaxy day

2 FROGS days

1.5 Statistics phyloseq day (under R)